$\gamma_{\Lambda}$ 

#### **REMARKS**

Claims 1-30 and 54 are pending in the application, and are subject to restriction. Independent claims 1, 20, and 54 have been revised to recite a solid oxide fuel cell that is prepared by a process whereby at least the solid electrolyte and anode being prepared by cofiring or sintering at least two layers to form a porous anode layer and a dense solid electrolyte layer, and then impregnating the porous anode layer with an aqueous solution containing a salt of at least ceria. Support for the amendments can be found throughout the specification and examples, and specifically at page 12, lines 16 and 17 for claim 54, ("aqueous impregnation using a concentrated salt solution), and page 13, lines 6 and 7 for claims 1 and 20 ("[o]ther components including ceria oxide (CeO<sub>2</sub>), are preferably added using the same impregnation methods. No new matter is presented by the amendments, the amendments simplify issues for appeal, and the amendments place the claims in condition for allowance. Accordingly, applicants respectfully request entry thereof and reconsideration of claims 1-30 and 54 in light of the following remarks.

Initially, applicants wish to thank Examiner Yuan for the courtesies extended the undersigned during the telephonic interview on January 21, 2005. Applicants' representative discussed the amendments now made to the claims, and the prior art, including a new document that has not yet been cited, but that has been cited in corresponding foreign applications. No agreement was reached, although the Examiner encouraged applicants to file the amendments presented herewith.

Turning now to the Action, claim 54 is alleged to be directed to an invention that is independent or distinct from the invention originally claimed. Specifically, the Action alleges that "a solid oxide fuel cell comprising a solid electrolyte... a ceramic-metal composite anode containing at least copper and a cathode, ... is a distinct species from a solid oxide fuel cell comprising a solid electrolyte ... a ceramic-metal composite anode containing at least copper and a cathode as recited in the original claims." (Action, at page 1, emphasis added). Applicants respectfully disagree with this allegation.

The originally presented claims that have been examined on the merits did not recite any species — copper or ceria. Rather, the claims that were originally presented and examined only recited a solid oxide fuel cell comprising a solid electrolyte comprised of an electronic insulator that allows transfer of anions, a ceramic-metal composite anode and a cathode. It was not until claim 54 was added that the species "ceria" was added to claims 1

and 20. Accordingly, a genus claim covering all of the species now recited in independent claims 1, 20, and 54 has already been examined. Applicants respectfully request that the Examiner reconsider and withdraw the restriction requirement.

Applicants appreciate with thanks the Examiner's withdrawal of the previous rejections under 35 U.S.C. §112, second paragraph.

On pages 2-6 of the Action, claims 1-6, 9-27, and 30 are rejected under 35 U.S.C. §103(a) as being unpatentable over Wallin, U.S. Patent No. 6,017,647 ("Wallin") in view of Anumakonda *et al.*, U.S. Patent No. 6,221,280 ("Anumakonda"). The present Action repeats the previous rejection. For the reasons set forth in applicants' responses filed on September 17, 2003, and August 11, 2004, this rejection is respectfully traversed.

Even assuming that Anumakonda suggests directing a sulfur-containing fuel to a solid oxide fuel cell, Anumakonda clearly states that the fuel cell must be "sulfur tolerant," or alternatively, have a provision for desulfurizing the product gas stream. See, col. 11, lines 58-63 of Anumakonda. The fuel cell of Wallin is not "sulfur tolerant," nor does Wallin include a provision for desulfurizing the product gas stream. Rather, Wallin's fuel cell is made from a conventional anode material, nickel (see, all of the examples). Applicants attach as an Appendix a series of publications that describe, inter alia, the poisoning effect of sulfur-containing impurities on an SOFC anode containing nickel or other conventional anode metals.

All of the solid oxide fuel cells prepared in Wallin's examples include an anode made from nickel — the examples are the only place in Wallin where preparation of an anode is described, since the remaining disclosure of Wallin pertains to the oxygen electrode, or cathode. The publications attached hereto reveal that it was known in the art at the time the present invention was made that sulfur-containing fuels poisoned or otherwise destroyed a fuel cell's operability when the fuel cell anode comprised nickel. Thus, combining Anumakonda with Wallin would have essentially destroyed the viability of Wallin's fuel cell. A person skilled in the art therefore would not have been motivated to combine Anumakonda and Wallin in the manner asserted in the Action.

Applicants traverse this rejection for the following additional reasons. Wallin discloses a composite oxygen electrode/electrolyte structure that contains a variety of materials. The oxygen electrode is the cathode. All of the examples of Wallin that

specifically disclose an anode material utilize nickel, and prepare the anode in a manner similar to the manner disclosed in some of the publications attached hereto. Applicants appreciate the one passing reference in Wallin (as graciously pointed out to applicants' representative by the Examiner during the interview) that the electrode/electrolyte can be the anode or cathode (col. 5, lines 33-36). Applicants respectfully submit that this sentence would not have suggested to the skilled artisan that modifying the anode of Wallin's examples to include cathode materials could somehow render Wallin's solid oxide fuel cell sulfur tolerant. Indeed, this one sentence in Wallin might possibly suggest, at best, that a nickel-containing anode, and an anode comprised of one of the oxygen electrode/electrolyte materials disclosed therein, would provide substantially equivalent results (*i.e.*, they are interchangeable). The present inventors discovered that this was not the case.

The present inventors discovered that anodes containing at least copper and/or ceria are capable of directly processing sulfur-containing fuels. The inventors discovered a method of incorporating relatively low melting materials (like copper) in an anode by initially co-firing the anode layer and the electrolyte layer, impregnating the resulting anode layer with a solution containing a salt of the anode material, and then heating the impregnated anode to drive off the liquid. Conventional methods of preparing anodes containing nickel (nickel cermet was combined with YSZ and co-fired together with the electrolyte layer at temperatures at or about 1250°C as disclosed in Wallin) were not suitable for preparing an anode containing copper since the copper would melt and diffuse out of the ceramic material during sintering. The inventors discovered that that anode needed to be sintered first, and then impregnated with a solution containing a salt of the anode material (e.g., Cu or CeO<sub>2</sub>). It was by this method that the inventors were able to prepare a solid oxide fuel cell capable of directly oxidizing sulfur-containing fuels.

Neither Anumakonda nor Wallin disclose making an anode in the manner recited in the claims. Wallin does disclose infiltrating the sintered fuel cell with an aqueous solution containing an electrocatalyst (*see*, paragraph bridging columns 5 and 6 and the examples); none of the electrocatalyst materials contain ceria or copper. The method of making the solid oxide fuel cell recited in the present claims therefore produces a novel and unobvious solid oxide fuel cell capable of directly oxidizing sulfur-containing hydrocarbons. The

Applicants do note that the paragraph bridging columns 3 and 4 disclose a laundry list of suitable "electronically-conductive materials" and "electrocatalysts," none of these materials includes ceria or copper.

On pages 6-9 of the Action, claims 1, 2, 7, 8, 20, 28 and 29 are rejected under 35 U.S.C. §103(a) as being unpatentable over Wallin in view of Fasano, *et al.*, U.S. Patent No. 6,051,330 ("Fasano"). The Action repeats the previous rejection. For the reasons set forth in applicants' response filed on September 17, 2003, and August 11, 2004, this rejection is respectfully traversed.

The Action apparently takes the position that the alcohol feeds disclosed in Fasano inherently have sulfur contents of from about 1 ppm to about 5000 ppm. The Examiner correctly notes the legal requirements to establish inherency insofar as the missing feature must necessarily be present in the prior art. The Action also states, however, that "both Fasano et al. and the present application utilize similar alcohols" (page 7 of the Action). The Action states further that "[t]he instant disclosure provides no further description with respect to the nature and characteristic of the alcohols used" (Action, at page 10). These are not accurate statements.

While it certainly is true that the present specification lists some of the same alcohols by name that are listed in Fasano, the present specification also makes clear that the alcohols disclosed and claimed therein contain at least 1 ppm sulfur. This is a very important difference between the alcohols recited in the present claims and the alcohols broadly disclosed in Fasano. As stated in previous responses, and shown by reference to a number of publications, the sulfur content of commercially available methanol for use in fuel cells is typically zero, and at most about 0.5 ppm. Absent any disclosure to the contrary, a person skilled in the art would recognize that the methanol disclosed in Fasano is precisely the type of methanol typically used in fuel cells — that is, methanol with no sulfur.

In sharp contrast to the Fasano disclosure, the alcohols described and claimed in the present specification require sulfur. That is, the present specification contains disclosure that is contrary to conventional wisdom with respect to fuel cells, which is that the alcohols and other fuels contain sulfur. To the extent that commercially available hydrocarbon fuels (such as an alcohol as recited in the dependent claims) contain no sulfur, sulfur can be added. The Examiner will note that sulfur is added to conventional decane in the examples to provide a

sulfur-containing fuel (thiophene was added to hydrocarbons to provide a sulfur-containing hydrocarbon). Accordingly, the combination of Fasano and Wallin would not result in a fuel cell containing a fuel comprising a sulfur-containing hydrocarbon, as recited in the present claims.

The combination of Fasano and Wallin also fails to suggest a fuel cell comprised of an anode containing at least ceria and/or copper, or the method of making the solid oxide fuel cell. Both Wallin and Fasano disclose conventional solid oxide fuel cell fabrication techniques (Wallin is discussed above, and Fasano, at col. 6, lines 7-33). The combination of Wallin and Fasano therefore would not result in the presently claimed solid oxide fuel cell. Applicants respectfully request that the Examiner reconsider and withdraw this rejection.

Applicants wish to bring to the Examiner's attention Cable, *et al.*, U.S. Patent No. 5,589,285 ("Cable"). Cable has been cited against claims similar to the originally presented claims (not the currently amended claims) in corresponding foreign applications. Cable was cited in applicants' Information Disclosure Statement dated June 3, 2003, and has been initialed by the Examiner. Accordingly, Cable is of record in the present application.

Cable appears to disclose sulfur tolerant fuel cells. The amended claims recite forming the electrolyte and anode by co-firing the two layers together, which is described in Cable as a "bonded" fuel cell. Cable teaches that bonded fuel cells could not process sulfur, and its disclosure therefore is directed to non-bonded fuel cells with various separator and microslip layers positioned between the electrolyte and electrodes. The present inventors therefore achieved what Cable said was not possible — a bonded sulfur-tolerant fuel cell.

In view of the foregoing, applicants respectfully submit that the present claims are in. condition for allowance. An early notice to this effect is earnestly solicited. Should there be any questions concerning this response, Examiner Yuan is invited to contact the undersigned at the telephone number listed below.

Respectfully submitted,

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#### Thermodynamics of sulfur chemisorption on metals. III. Iron and cobalta)

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Isosteric heats of chemisorption of sulfur on iron and cobalt have been evaluated from measurements of the equilibrium ratio of H<sub>2</sub>S/H<sub>2</sub> gas mixtures in contact with the powdered metals over a temperature range from 677 to 855 K. At low surface coverage of sulfur on cobalt, the heat of chemisorption with respect to 1/2 S2(g) is -236±15 kJ mol<sup>-1</sup>. With increasing coverage it decreases to -200±10 kJ mol<sup>-1</sup>. The heat of sulfur chemisorption on iron is  $-197 \pm 15$  kJ mol<sup>-1</sup>. For equivalent coverage, the thermodynamic activity of sulfur chemisorbed on cobalt exceeds that of iron by approximately one order of magnitude. At low surface coverage (<0.5 saturation coverage), the bond strength of sulfur on the various metals is found to vary in the order: Ni > Co > Ru > Fe.

#### INTRODUCTION

Sulfur dissolved in metallic iron has a very large free energy potential for segregation to the surface. Preferential surface or grain boundary segregation of small quantities of dissolved sulfur can have profound effects on the mechanical and corrosive properties of iron and its alloys. 1 Poisoning by trace concentrations of gaseous sulfur compounds can alter selectivity and severely limit the lifetime of iron-based catalysts used in such important reactions as the Fischer-Tropsch synthesis<sup>2</sup> and ammonia synthesis. 3 Likewise cobalt and iron-cobalt alloys are susceptible to deleterious catalyst poisoning. The purpose of the present work is to examine the thermodynamics of sulfur chemisorption on iron and cobalt as part of a series of equilibrium sulfur chemisorption studies on various metal surfaces. Previous measurements have dealt with nickel4 and ruthenium.5

Indirectly the thermodynamics of reversible sulfur chemisorption on polycrystalline iron at 1123 K were evaluated by Grabke et al. 6-8 by measuring the changes in the initial rates of nitrogenation (with N<sub>2</sub>) and carburization (with CH<sub>4</sub>) following equilibrium exposure to H<sub>2</sub>S-H<sub>2</sub> gas mixtures. Paulitschke et al. 7-9 estimated the heat of formation of adsorbed sulfur from segregation studies on iron foils at 973 to 1373 K, using Auger electron spectroscopy to measure the sulfur surface coverage in equilibrium with dissolved sulfur in the bulk. Agrawal et al. 10 have shown that the cobalt-catalyzed methanation reaction is almost completely poisoned by as little as 15 ppb H<sub>2</sub>S in H<sub>2</sub> at 663 K. It is obvious that the surface coverage with sulfur approaches saturation at sulfur chemical potentials well below those required to produce the bulk sulfides of iron and cobalt.

#### **EXPERIMENTAL**

The iron and cobalt samples used in this study were prepared by precipitation of 0.5 M aqueous solutions of ferric or cobaltous nitrate with 60% excess potassium carbonate at 298 K. The precipitates were washed

thoroughly with distilled water, dried to 373 K, reduced stepwise in a hydrogen stream at 0.1 MPa for periods of 2 h each at 423, 573, and 723 K, and subsequently cooled to 298 K in helium. The specific surface areas of the two powders were measured by CO adsorption at 298 K following reduction in hydrogen at 773 K for 2 h. For iron the saturation coverage was found to correspond to 20.7  $\mu$ mol CO/g, and for Co, 24.9  $\mu$ mol CO/g. The specific surface areas were calculated to be 1.17  $m^2/g$  of Fe and 1.40  $m^2/g$  of cobalt, assuming 1.1×10<sup>15</sup> CO atoms adsorbed per cm<sup>2</sup> of metal surface area.

The experimental techniques used in this study have been described previously. 4,5 Briefly, we place a sample of the metallic powder ( $\sim 5 \times 10^{-3}$  g) on the quartz fritted disk of a microreactor. The microreactor is part of a quartz/Teflon closed loop through which hydrogen recirculates. Sampling valves are used to inject aliquots of a dilute H2S in H2 mixture of known composition into the loop and to take samples of the recirculating gas for analysis of the H<sub>2</sub>S/H<sub>2</sub> ratio. For H<sub>2</sub>S analysis a gas chromatograph is used with a very sensitive photoionization detector, so that H2S concentrations below 1 ppb in H<sub>2</sub> can be detected. After uniform sulfur adsorption coverage is established, isosteres are determined by varying the reactor temperature and measuring the changes in the H<sub>2</sub>S/H<sub>2</sub> ratio of the recirculating gas. Because the amount of sulfur adsorbed is several orders of magnitude greater than the amount of H2S in the recirculating gas, transfer of sulfur from the surface to the gas phase is sufficiently small during the measurements to have an insignificant effect on surface coverage.

#### **RESULTS**

#### iron

Isosteres for sulfur chemisorption on iron were determined from the measured thermodynamic chemical potential of gaseous sulfur (as H,S in H, at a total pressure of 1 atm) for several values of sulfur coverage (Fig. 1). The enthalpy and entropy for reversible sulfur adsorption [S(a)]

 $H_2S(g) = S(a) + H_2(g)$ ,

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<sup>&</sup>lt;sup>a</sup>Support of the Office of Basic Energy Sciences, Division of Chemical Sciences of the U.S. Department of Energy is gratefully acknowledged.

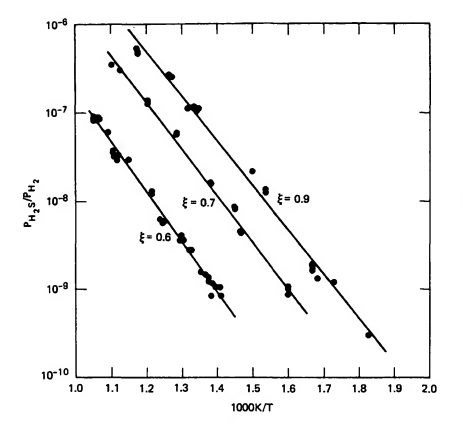


FIG. 1. Sulfur chemisorption isosteres on iron powder.

were calculated from a Van't Hoff analysis of the isosteric data (Table I). Standard enthalpies and entropies of formation for adsorbed sulfur [S(a)] (Table II) were evaluated from the isosteric adsorption data and the tabulated thermodynamic properties of  $H_2S$  and  $H_2$  with  $S_2(g)$  as the reference state  $H_2S$  using the relationships

$$\Delta H_f^{\circ}(T)[S(a)] = \Delta H_{ads}(T) + \Delta H_f^{\circ}(T)[H_2S(g)],$$

and

$$S_f^{\circ}(T)[S(a)] = \Delta S_{ads}(T) + S_f^{\circ}(T)[H_2S(g)] - S_f^{\circ}(T)[H_2(g)] - \frac{1}{2}S_f^{\circ}(T)[S_2(g)],$$

where  $\Delta H_{f}^{o}(T)$  and  $S_{f}^{o}(T)$  represent the standard enthalpy and entropy of formation at temperature T. The mean experimental temperature of each isostere was chosen as the reference temperature in the calculations. The

normalized sulfur coverage  $\xi$  represents the amount of sulfur introduced into the system divided by the CO adsorption capacity at 298 K of the iron powder.

The enthalpies and entropies of sulfur adsorbed on iron are qualitatively similar to those observed previously for nickel<sup>4</sup> and ruthenium.<sup>5</sup> First, the sulfur adspecies are atomic sulfur since adsorption as HS(a) or  $H_2S(a)$  would be associated with a much larger entropy of formation. Second, the enthalpy decreases substantially with increasing sulfur coverage.

#### Cobalt

The chemisorption isosteres of sulfur on cobalt (Fig. 2) were obtained at considerably lower surface coverages -normalized to the CO adsorption capacity of the

TABLE I. Analysis of isosteres for sulfur chemisorption on iron and cobalt.

Metal	Coverage (ξ) <sup>2</sup>	ΔH <sub>ads</sub> (kJ mol <sup>-1</sup> )	ΔS <sub>ads</sub> (J mol <sup>-1</sup> K <sup>-1</sup> )	T <sub>arg</sub> (K)	$n^{\mathfrak{b}}$	rc
Iron	0.6	-107.8 ± 9.2	21.9±11.1	826	37	0.996
	0.7	$-100.7 \pm 8.8$	$10.9 \pm 12.2$	725	21	0.996
	0.9	$-96.5 \pm 10.4$	$4.9 \pm 15.4$	677	22	0.994
Cobalt	0.09	$-145.1 \pm 18.4$	$2.9 \pm 21.5$	855	36	0.992
	0.18	$-143.4 \pm 9.4$	$-0.4 \pm 11.1$	845	14	0.998
	0.3	$-150.8 \pm 8.3$	$-14.5 \pm 10.0$	828	15	0.998
	0.4	$-126.7 \pm 7.4$	$9.2 \pm 8.8$	844	4	0.998
	1.0	$-110.2 \pm 4.4$	$-13.3 \pm 6.2$	702	9	0.999

 $<sup>^{</sup>a}\xi$  = coverage normalized to CO uptake at 298 K.

 $b_n$  = number of data points in isostere.

 $<sup>^{</sup>c}r = least-squares$  correlation coefficient.

TABLE II. Isosteric heat and entropy of formation of sulfur chemisorbed on iron and cobalt<sup>b</sup>

Metal	Coverage (ξ) <sup>a</sup>	ΔH <sup>o</sup> <sub>f</sub> (kJ mol <sup>-1</sup> )	S°(J mol <sup>-1</sup> K <sup>-1</sup> )	Temperature range (K)
Iron	0.6	-197	105	954-711
	0.7	-189	95	911-625
	0.9	<b>- 185</b>	86	855-547
Cobalt	0.09	- 234	87	901-803
	0.18	- 233	84	898-796
	0.3	- 240	69	903-766
	0.4	-216	93	901-794
	1.0	- 199	68	752-611

 $<sup>{}^{8}\</sup>xi$  = sulfur coverage normalized to CO uptake at 298 K.  ${}^{9}Reference$  state =  ${}^{1}_{2}S_{2}(g)$ .

sample). The enthalpy of sulfur adsorption on cobalt is found to be greater than on iron (Table I). Again the standard entropy of adsorption for low surface coverage (Table II) suggests S(a) as the adspecies rather than HS(a) or  $H_2S(a)$ . At low surface coverage the magnitude of the standard enthalpy of formation of chemisorbed sulfur is approximately 38 kJ mol<sup>-1</sup> greater for cobalt

than for iron. However, as the sulfur coverage increases,  $\Delta H_{\rho}^{\rho}$  for chemisorption on iron and cobalt (Fig. 3) approaches the values for  $\frac{1}{8}$  Co<sub>9</sub>S<sub>8</sub> and FeS, respectively. Also the enthalpy difference between chemisorbed sulfur on Fe and Co approaches the difference for the respective bulk sulfides. <sup>12,13</sup>

#### DISCUSSION

In comparing our results with published thermodynamic values we find that our heat of formation of chemisorbed sulfur on iron at low surface coverage  $(\Delta H_f^o[S_a]) = -197 \text{ kJ mol}^{-1}$ , Table I) is very close to that calculated from the sum of the heat of segregation<sup>9</sup> (-165 kJ mol<sup>-1</sup>) and dissolution<sup>14</sup> (-31 kJ mol<sup>-1</sup>) for polycrystalline iron. Assuming that the saturation coverage of sulfur equals the uptake of carbon monoxide, we find by extrapolation of our data that the equilibrium sulfur activity in the gas  $(H_2S/H_2)$  for one-half saturation coverage at 1123 K corresponds to 0.7 ppm. From the data by Grabke et al. <sup>8</sup> one obtains a sulfur activity of 0.8 ppm  $(H_2S/H_2)$  for this degree of surface coverage at 1123 K. The agreement is most satisfactory considering the different experimen-

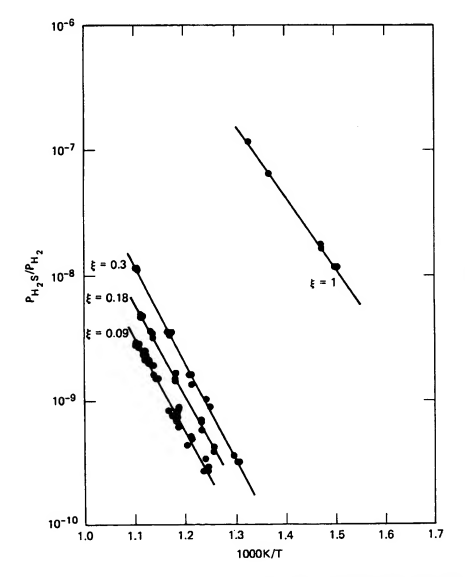


FIG. 2. Sulfur chemisorption isosteres on powdered cobalt.

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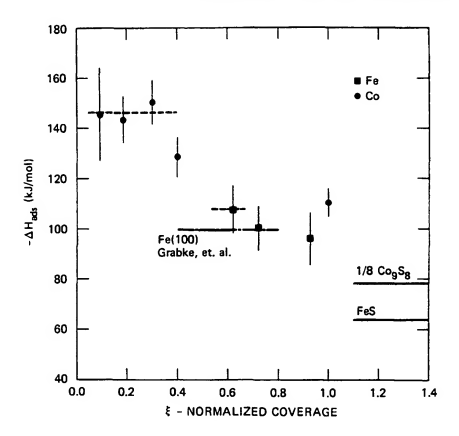


FIG. 3. Heat of sulfur chemisorption vs coverage on Fe and Co powders.

tal techniques employed. The data show that even at very high temperatures the thermodynamic activity of sulfur is extremely low for fractional coverage with chemisorbed sulfur.

It is valuable to examine the entropy of formation of chemisorbed sulfur  $S_j^*(T)[S(a)]$ . In principle this parameter can be estimated from an analysis of the vibrational modes of the adsorbed phase. However, vibrational data for sulfur adsorbed on all of the metals studied are unavailable; in addition, one needs to consider other contributions to the entropy such as adatom-substrate interactions and the configurational contributions. Nevertheless, sufficient data exist to estimate the standard entropy of formation of chemisorbed sulfur on a metal, such as nickel, and to compare it with the entropies of formation measured for Fe, Co, and Ru.

The higher frequency-stretching modes<sup>15</sup> for adsorbed sulfur on Ni(100) are found to be 46 meV for the  $p(2\times2)$ structure, and 44 meV for the  $C(2\times2)$  structure. We approximate the collective vibrational modes of sulfur chemisorbed [as  $C(2\times2)$ ] on nickel (100) by vibrations with the frequencies 355, 258, and 135 cm<sup>-1</sup>, corresponding to high resolution electron energy loss spectroscopy (EELS) peaks at 44, 32, and 16.7 meV, respectively. although we recognize that the actual surface phonon frequencies are distributed. 16 Similar vibrational modes were found for sulfur chemisorption on Ni(111). 17 Treating each mode as a one-dimensional harmonic oscillator, 18 i.e., using the Einstein model, we have calculated the vibration entropy for sulfur adatoms on Ni at 850 K and obtained a value of 48.5 J mol-1 K-1. Considering the uncertainty of the atom frequency distribution, the probable variation in frequency distribution with surface structure, and the unknown changes in the substrate surface vibrational modes due to the presence of the adsorbed phase (for example, oxygen adsorption of Ni may increase the vibrational frequency of the nickel atoms at the surface), <sup>19</sup> we estimate that the actual vibration entropy of sulfur chemisorbed on Ni is within the range  $50\pm20~\mathrm{J}~\mathrm{mol}^{-1}~\mathrm{K}^{-1}$  for temperatures from 800 to 900 K. This value is somewhat less than the experimental entropies of formation for sulfur adsorbed on Ni (Table III), but is comparable to the entropies for sulfur in bulk sulfide, taken to be the difference between the standard entropies of formation of the bulk sulfide and the pure metal. i.e.,

$$S_f^{\circ}(T)[\text{sulfide}] = S_f^{\circ}(T)[\text{Me}_xS] - xS_f^{\circ}(T)[\text{Me}]$$
.

Similar considerations apply to cobalt. However, for chemisorbed sulfur on Fe and Ru<sup>20</sup> the experimental entropies are significantly greater by about 30 J mol<sup>-1</sup> K<sup>-1</sup>.

The configurational contribution to the entropy of adsorption is determined by the sulfur surface coverage and the nature of the adsorption. For noninteracting adatoms with a single binding energy, the Langmuir adsorption isotherm applies, and

$$S_f^{\circ}(\text{config}) = -R \ln[\theta/(1-\theta)]$$
.

For repulsive nearest neighbor adatom interactions, at low coverage,  $S_{\rho}^{\rho}(\text{config}) \cong -R \ln \theta$ . But as the coverage approached one-half monolayer [for a (100) surface], the configurational entropy would decline sharply until all noninteracting sites were filled. The entropy would then suddenly increase as the nearest neighbor sites started

TABLE III. Thermodynamics of sulfur chemisorption of metals.

			$S_f^{\circ}(T_{avg})[S(a)]^b$	(J mol <sup>-1</sup> K <sup>-1</sup> )	$\Delta H_f^{\circ}(T_{avg})[S(a)]$	(kJ mol <sup>-1</sup> K <sup>-1</sup> )
Metal	Coverage (ξ) <sup>a</sup>	$T_{ayg}({ m K})$	Experimental	Calculated	Experimental	Calculated
Ni	0.63	863	71	49	- 249 + 17	- 268
	0.72	826	52	48	$-248 \pm 12$	- 251
	0.75	806	47	47	$-244 \pm 18$	- 244
	0.92	701	52	43	- 216 + 5	- 222
	0.98	625	66	41	$-196 \pm 26$	- 212
	1.12	536	72	37	-174 + 8	- 193
Co	0.09	855	68°	55	- 234 ± 18	- 246
	0.18	845	69 <sup>c</sup>	55	-233 ± 9	- 245
	0.3	828	62 <sup>e</sup>	54	- 240 ± 8	- 247
	0.4	844	89°	55	$-216 \pm 7$	- 245
	1.0	702	68	51	$-199 \pm 4$	-211
Fe	0.6	826	105	74	- 197 ± 9	- 223
	0.7	725	95	72	$-189 \pm 9$	- 206
	0.9	677	86	70	$-185 \pm 10$	- 196
Ru	0.67	886	97	48	- 215 ± 39	- 258
	0.75	818	83	46	$-213 \pm 9$	- 243
	0.83	776	53	45	$-216 \pm 20$	- 222
	1.00	510	22	37	- 187 ± 9	-179

<sup>&</sup>lt;sup>a</sup>ξ: coverage normalized to CO uptake at 298 K.

S<sub>f</sub> for sulfur adsorption on cobalt has been adjusted by  $+R \ln[\xi(1-\xi)]$  for  $\xi \le 0.4$  to allow for configurational entropy.

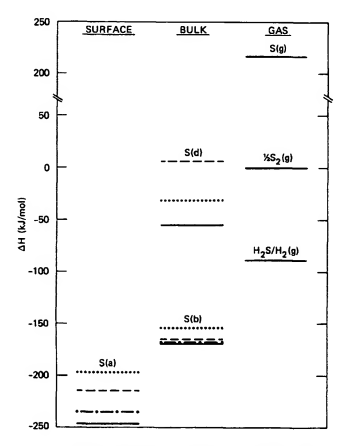


FIG. 4. Energetics of sulfur interaction with metals relative to  $\frac{1}{2} \; S_2(g)$  .

filling (at higher energy), declining again as monolayer saturation coverage was reached. For two-dimensional phase transformations, as might be expected for very attractive nearest neighbor interactions, the configurational term would vary as  $-\ln\theta$  up to the nucleation point and remain nearly constant thereafter.

We conclude that the configurational entropy term for sulfur chemisorption should vary approximately as  $R \ln \phi$  for  $\phi \ll 0.5$  (where  $\phi$  is the surface coverage with sulfur relative to saturation coverage). For  $\phi \geq 0.5$ , the S<sub>f</sub>(config) term is very difficult to predict since the isotherms obviously do not follow the Langmuir model. However, we were able to measure the isosteres for cobalt with sulfur coverages to less than one-tenth of the CO adsorption capacity; therefore in Table III, the Langmuir configurational entropy term was added to the estimated entropy of formation of adsorbed sulfur.

The measured magnitude of the heats of adsorption of sulfur at low coverage on the metal surfaces examined thus far follow the order Ni> Co> Ru> Fe. The free energy ( $\Delta G^{\circ}$ ) of sulfur adsorption at low coverage on these metals at 850 K follows the same order, Ni> Co> Ru> Fe, although the difference between the  $\Delta G^{\circ}$ , values for Ni, Co, and Ru is much less than that between the values of  $\Delta H^{\circ}_f$ . Since essentially the same degrees of freedom are lost during adsorption and during formation of bulk compounds, one can suggest that  $S^{\circ}_f$  for adsorption should be approximately the same as  $S^{\circ}_f$  for bulk compound formation. If the entropy of adsorbed sulfur were taken equivalent to the entropy of the sulfides and the values for  $\Delta H^{\circ}_f$  were adjusted to keep  $\Delta G^{\circ}_f$  constant at

References for bulk sulfide data are: iron and ruthenium, Ref. 11; nickel and cobalt, Ref. 21. For ruthenium, S\*(298)[RuS2(8)] was taken as 14.5 cal mol<sup>-1</sup> K<sup>-1</sup>.

 $T_{\rm avg}$  (Table III), the heats of formation for chemisorbed sulfur would be  $-254~{\rm kJ\,mol^{-1}}$  for Ni,  $-246~{\rm kJ\,mol^{-1}}$  for Co,  $-241~{\rm kJ\,mol^{-1}}$  for Ru, and  $-215~{\rm kJ\,mol^{-1}}$  for Fe. Again, the same order prevails, although the magnitudes of  $\Delta H_{\rm f}^{\rm o}$  would be increased for each metal.

The chemisorption of sulfur on these four metals exhibit more similarities than differences. The experimental heats of formation of sulfur adatoms (at 800 K for low sulfur coverage) are considerably below the corresponding levels for the bulk sulfides (see Fig. 4). The ΔH, levels for the bulk sulfides follow the same relative order as adsorbed sulfur, although the differences between Ni, Co, and Ru may not be significant for both adsorbed sulfur and bulk sulfides. The experimental entropy of adsorbed sulfur equals or exceeds that of the most stable bulk sulfide. The very large heat of segregation of dissolved sulfur to the surfaces of iron, cobalt, and nickel, in addition to the favorable entropy change, provides an energy potential of (135±30 kJ mol<sup>-1</sup>) for the well-known tendency of dissolved sulfur to segregate to the surfaces of these metals. (Presumably this trend applies to ruthenium as well, although the data for sulfur dissolved in Ru are lacking). The bond strength of a sulfur adatom at 800 K, i.e., the heat of adsorption from a monatomic gas, shows some variation from metal to metal, ranging from 464 kJ mol-1 for nickel to 414 kJ mol-1 for iron.

The present results are in good agreement with the correlation of Benard et al. <sup>23</sup> linking the heats of formation of two-dimensional metal-sulfur surface species to the heats of formation of the bulk sulfides. If we presume that the two-dimensional sulfide corresponds to the low coverage ( $\xi \le 0.7$ ) results for Ni, Fe, Co, and Ru, then our data for  $\Delta H_{f}^{o}$  (800 K) [S(a)] are close to the proposed trend line, which follows approximately the relation  $\Delta H_{f}^{o}[S(a)]_{2D} \cong 1.23 \Delta H_{f}^{o}[(1/y) \text{Me}_{x} \text{S}_{y}]$ , indicating that the heat of formation of chemisorbed sulfur on a metal

surface is 20% to 25% greater than the equivalent heat of formation of the bulk sulfide of the metal.

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### Thermodynamics of sulfur chemisorption on metals. I. Alumina-supported nickel

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Sulfur chemisorption isosteres have been measured for nickel in powdered form and for nickel supported on two different alumina powders. The experiments were conducted in a closed-loop gas recirculation system containing one atmosphere hydrogen. Isotherms were determined by stepwise injection of H<sub>2</sub>S aliquots into recirculating hydrogen gas and analyzing for the H<sub>2</sub>S concentration as equilibrium was approached. Isosteres were measured by varying the sample temperature and monitoring the H<sub>2</sub>S/H<sub>2</sub> ratio in the gaseous environment. A gas chromatograph and a photoionization detector was used to measure the H<sub>2</sub>S concentration to levels below 1 ppb. As monolayer coverage is approached the H<sub>2</sub>S/H<sub>2</sub> ratio attains the equilibrium values reported for the bulk sulfide, Ni<sub>3</sub>S<sub>2</sub>. Adsorbed sulfur is very strongly bound to the surface of nickel. The heat of formation of chemisorbed sulfur with respect to 1/2 S<sub>2</sub>(g) at 800 K is 247 kJ mol<sup>-1</sup> more negative than the heat of formation of Ni<sub>3</sub>S<sub>2</sub>. The heat of segregation exceeds 190 kJ mol<sup>-1</sup>. The results demonstrate that the equilibrium sulfur coverage exceeds one half monolayer on Ni/Al<sub>2</sub>O<sub>3</sub> unless the H<sub>2</sub>S/H<sub>2</sub> ratio is less than 1 ppb even at temperatures as high as 940 K.

#### I. INTRODUCTION

Nickel catalysts, presently the most suitable material for the efficient production of methane from coal-derived synthesis gas, are highly susceptible to deactivation by trace contaminants of sulfur compounds in the feedstocks. Although deactivation of nickel methanation catalysts has been observed with as little as 0.3 ppm H<sub>2</sub>S in the inlet gas, <sup>1</sup> the thermodynamic aspects of sulfur poisoning on nickel have not been determined with sufficient accuracy to establish the threshold levels of sulfur contaminants above which the catalyst surface accumulates a specified fractional sulfur coverage. The purpose of the present study is to provide basic thermodynamic information relating the surface coverage of sulfur on metallic nickel crystallites to the gas phase sulfur activity in a reducing atmosphere.

The reversible adsorption of sulfur on nickel surfaces was first reported by Perdereau and Oudar<sup>2</sup> in 1970 for a polycrystalline foil, followed in 1971 by Rostrup-Nielsen<sup>3</sup> for nickel supported on MgO · Al<sub>2</sub>O<sub>3</sub>. Both authors measured sulfur chemisorption isotherms with sulfur activity (defined by the ratio of H<sub>2</sub>S to H<sub>2</sub>) of the order  $10^{-6}$  at elevated temperatures. These studies suggested that the free energy of formation of chemisorbed sulfur was at least 50 kJ mol<sup>-1</sup> more negative than the most stable bulk sulfide Ni<sub>3</sub>S<sub>2</sub>. A Recently, Oliphant *et al*. published H<sub>2</sub>S desorption isotherms for several supported nickel catalysts. An enthalpy of adsorption of -160 kJ mol<sup>-1</sup> and a free energy of formation -45 kJ mol<sup>-1</sup> of chemisorbed sulfur were reported.

The present study extends the measurement of sulfur adsorption isotherms and isosteres on supported nickel surfaces to considerably lower levels of sulfur activity than previously attained. By means of a specially designed system incorporating a higher sensitivity sulfur detection device, we were able to determine reliably the equilibrium concentration of  $\rm H_2S$  in one atmosphere of  $\rm H_2$  at levels below 1 ppb.

We have also examined the chemisorption of sulfur on the (100) and (111) planes of single crystal nickel using a technique similar to that of Perdereau and Oudar. The results of the study on single crystal nickel will be discussed in detail in a subsequent paper. <sup>6</sup>

#### II. EXPERIMENTAL DETAILS

#### System design and operation

A closed-loop gas-recirculation system has been assembled and used to measure isosteric heats of sulfur chemisorption on alumina-supported nickel crystallites and powdered nickel samples. Figure 1 is a schematic diagram of the apparatus. The heart of the apparatus is a Teflon and quartz loop, represented in Fig. 1 as a bold line. Essential equipment in the closed loop include an electrically heated quartz microreactor which retains the catalyst sample; two Teflon rotary valves, one for gas analysis (I2), the other for injecting sulfur (I3); and a Teflon diaphragm pump with a glass bulb to dampen pressure fluctuations. The loop is isolated by Teflon diaphragm valves (V8 and V9). During operation H<sub>2</sub>S is

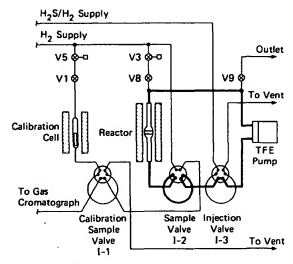


FIG. 1. Sulfur chemisorption system for dispersed metal catalysts.

injected into the loop and the recirculating gas is analyzed for  $H_2S$  by a special gas chromatograph column and detector.

The lower sensitivity limit of the gas chromatographic detection system is 0.1 ppb  $\rm H_2S$ . The sensitivity is limited by baseline drift rather than noise sources. The column is a 6-ft. length of FEP tubing with 0.095 in. i.d. packed with Chromosil 310 (a treated silica gel designed to separate  $\rm H_2S$ ). The detector is a photoionization cell, HNU Model PI-51, equipped with a 10.2 eV hydrogen lamp. Both detector and column are operated at 298 K in a hydrogen carrier stream of 30-40 cm³/s. The sampling volume is 1.26 cm³ (12).

For detector calibration, a solid-state electrochemical cell Ag<sub>2</sub>S/AgI/Ag, generates known concentrations of H<sub>2</sub>S in a hydrogen stream. The nonstoichiometries of the  $\alpha$  and  $\beta$  forms of Ag<sub>2</sub>S, first observed by Wagner, <sup>7</sup> were verified by monitoring the cell voltage and current as a function of time with the cell under vacuum. The rate of H2S generation was limited by the kinetics of reaction of  $H_2$  with  $\beta$ -Ag<sub>2</sub>S at 410 K. Increasing the cell voltage had little effect on the rate. Application of 25 to 50 mV potential between the electrodes allowed Ag\* ions to migrate to the Ag electrode and generate a measured electric current in the cell. The H2S generation rate was equal to one-half the cell current inasmuch as electric leakage across the AgI ionic conductor was small at 50 mV. The cell was reliably accurate for calibration at 0.1 ppm and above.

Adsorption isotherms for the nickel-sulfur system are determined by injecting  $H_2S$  aliquots stepwise into the recirculation loop and analyzing the recirculating gas for  $H_2S$  after equilibrium is reached. After a uniform sulfur coverage is attained at the highest temperature examined for that coverage, precise isosteres are measured below 1 ppm by decreasing then increasing the catalyst temperature and monitoring the  $H_2S/H_2$  ratio while the gas circulates over the catalyst. The coverage was considered uniform when the isosteric data points agreed for decreasing and increasing temperature. However, when the  $H_2S$  concentration exceeds 1 ppm, the recirculating gas (260 cm³) accumulates a significant amount of sulfur, thus lowering the average sulfur coverage.

The concentration of  $\rm H_2S$  in the recirculating gas stream can be analyzed via injection valve I2 and the gas-chromatograph column and detector. Thus, one can follow the approach to equilibrium as a function of time and thereby gain some indication of transport rates and adsorption kinetics. Because the sampling volume is small, depletion of  $\rm H_2S$  by aliquot sampling is insignificant.

The closed-loop system places severe demands on the materials of construction. The materials must (a) be inert to irreversible  $H_2S$  adsorption and reaction, (b) have minimal sites for reversible adsorption to avoid sinks and sources of  $H_2S$  with changes in pressure, and (c) seal leak tight to prevent air diffusion into the loop or  $H_2$  and  $H_2S$  leakage out of the loop. We decided to use a quartz reactor and to use tetrafluoroethylene polymer (TFE Teflon) and quartz as the materials of construction

whenever possible because  $\rm H_2S$  adsorbs on metal and most glasses at room temperature. § For sampling, sixport rotary Teflon valves (11, 12, 13) rigged with injection loops are satisfactory. The internal components of the recirculation loop were constructed entirely of quartz and Teflon. We did not observe adsorption of  $\rm H_2S$  when it was injected into the loop with the reactor empty. Heating the quartz bulb previously exposed to  $\rm H_2S$  from 30 to 80 °C desorbed only 0.3 nmol  $\rm H_2S$ . Thus, the loop was inert and had minimal  $\rm H_2S$  adsorption capacity for 1 ppm  $\rm H_2S$  in 1 atm  $\rm H_2$  at room temperature.

However, leakage or permeation in the valves and fittings allowed air to accumulate in the loop at a rate of 1 ppm/min. The  $\rm H_2S$  partial pressure was not noticeably affected by this accumulation of air. Most likely, the oxygen that leaked into the system was converted to water. The resulting  $\rm H_2O$  levels attained were estimated to be less than 100 ppm during measurement periods.

Interference in our measurements by atmospheric constituents at 1 ppm/min leak rates is minimal because conversion of  $H_2S$  or chemisorbed sulfur to  $SO_2$ , COS,  $CH_3SH$ ,  $CS_2$ , or sulfate ion are thermodynamically unfavorable in a reducing atmosphere of  $H_2$  at 723 K.

#### Catalyst preparation

Sulfur chemisorption isosteres have been measured for three nickel samples. The first was a powdered nickel sponge obtained from Johnson Matthey and Co. (spectroscopically pure). Based on CO adsorption at 300 K, the surface area of the powder was very low (0.26 m²/g) (assuming CO saturation coverage = 1.1  $\times 10^{15}$  molecule/cm²) after 120 min reduction in H<sub>2</sub> at 773 K. The sample continued to sinter at 773 K. After 16 h of reduction in 1 atm H<sub>2</sub>, the specific surface area had decreased to 0.20 m²/g.

The second sample was a 5 wt% Ni catalyst supported on alumina. The support, Condea Pural A, a high-surface-area (350 m²/g)  $\gamma$ -alumina, was calcined in pure  $O_2$  at 773 K for 60 min, and then impregnated with an aqueous solution of Ni(NO<sub>3</sub>)<sub>2</sub> by the incipient wetness technique. The catalyst was dried in air and calcined in  $O_2$  at 623 K for 100 min. For adsorption studies a sample (3.2 mg) was loaded into the reactor and reduced in situ in flowing  $H_2$  for 750 min at 825 K. Based on CO adsorption, we estimate a specific surface area of 75 m²/g nickel.

The third sample was Ni (5 wt%) supported on a low-surface-area alumina (20 m²/g 65%  $\alpha$ -alumina). The support was calcined in pure oxygen at 1100 K for 8 h, ground, screened to a mean particle size of  $80\pm20~\mu\text{m}$ , and then impregnated with Ni(NO)3 solution. The catalyst sample (18.2 mg) was reduced in situ in pure H2 for 15 h at 775 K and 5 h at 930 K. The specific metal surface area was 51 m²/g, based on CO adsorption at 310 K. Table I summarizes the properties of the three nickel samples used in this study.

#### III. EXPERIMENTAL RESULTS

The isosteric heats for sulfur chemisorption on three nickel catalysts were determined from the adsorption

TABLE I. Nickel catalyst properties.

			Specific surface	area* (m²/g)
Catalyst	Weight (mg)	Support	Support	Metal
Ni sponge	40	•••	•••	0, 20
5 wt δ Ni/γ-Al <sub>2</sub> O <sub>3</sub>	3, 2	$\gamma$ -Al <sub>2</sub> O <sub>3</sub>	350	75
5 wt δ Ni/γ-Al2O3	18. 2	65% α-Al <sub>2</sub> O <sub>3</sub>	20	51

 $^{2}$ Based on CO uptake at 300 K on exposure to 1 vol% CO in He and on the assumption of 1.1×10<sup>15</sup> CO adsorption sites per cm<sup>2</sup> of metal.

isosteres (Fig. 2). The parameter  $\xi$  represents the fractional surface coverage by sulfur relative to CO adsorption. At the sulfur coverages used in these studies, we calculate from these data the heats and entropies of adsorption (Table II), based on the equation

$$P_{\rm H_2B}/P_{\rm H_2} = K_{\rm p} = \exp(\Delta H/RT - \Delta S/R)$$
, (1)

where  $\Delta H$  and  $\Delta S$  are relative to gaseous  $H_2S$  and  $H_2$ . The surface coverage  $\xi$  for the Ni sponge isostere (Fig. 2) could not be precisely determined because of the uncertainty of the degree of sintering, but it probably equals or exceeds 0.5. Despite the differences in the nature of the support, the dispersion of the nickel crystallites, and the relative sulfur coverage, the enthalpies and entropies of adsorption are the same within experimental uncertainty for the three catalysts.

A series of isosteres with increasing sulfur coverage were determined for the Ni/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub> (5 wt %) catalyst (Fig. 3). The heats of adsorption calculated from the isosteres (Table III and Fig. 4) exhibit three regimes. At relatively low surface coverage of sulfur  $(\xi < 0.75)$ , the heat of adsorption has an average value of  $\langle \Delta H_a \rangle_{**}$ = -157 kJ mol<sup>-1</sup>. At higher coverage it slowly decreases and then exhibits a discontinuous increase for  $\xi > 1.12$ . The adsorption entropies remain nearly constant then increase discontinuously above  $\xi = 1.12$  (Table III), coincident with the rapid increase in  $\Delta H_a$ . The discontinuity in  $\Delta H_a$  and  $\Delta S_a$  probably represent a change in the adsorbing species, from S adatoms to HS(a) or  $H_2S(a)$ . At the lowest coverage,  $\xi = 0.63$ , for which isosteres were measurable on Ni/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub> (5 wt %), the sulfur activity was very low, reaching only 5 ppb at 932 K.

Adsorption isotherms were evaluated at 823, 623, and 473 K (Table IV) from the data represented in Fig. 3 for Ni/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub>. The equilibrium sulfur activity shows an exponential increase with increasing coverage e.g., eight orders of magnitude at 623 K. The isotherms can-

TABLE II. Sulfur adsorption equilibria on various nickel catalysts.

Catalyst	Coverage relative to CO uptake (ξ)	Adsorption enthalpy $\Delta H_{\rm ads}$ (kJ mol <sup>-1</sup> )	Adsorption entropy $\Delta S_{ads}$ J mol <sup>-1</sup> K <sup>-1</sup> )
Ni sponge	> 0. 5	-143.5±18.0	35.9
$Ni/\gamma - Al_2O_3$ (5.0 wt%)	0.64	-143.1±16.3	39.6
$Ni/\alpha-Al_2O_3$ (5.0 wt)	0.75	$-155.2 \pm 17.7$	35.9

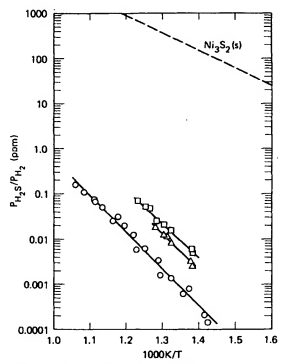


FIG. 2. Thermodynamic activity of sulfur chemisorbed on supported nickel. (o) 5 wt% Ni/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub>,  $\xi$  = 0.75; (n) 5 wt% Ni/ $\gamma$ -Al<sub>2</sub>O<sub>3</sub>,  $\xi$  = 0.64; (a) Ni sponge,  $\xi$  = 0.5, where  $\xi$  is the coverage normalized to CO uptake at 300 K.

not be interpreted in terms of Langmuir adsorption, i.e., sulfur adatoms bound to localized noninteracting sites.

The rate of approach to equilibrium was generally slow even at temperatures as high as 900 K. After H<sub>2</sub>S

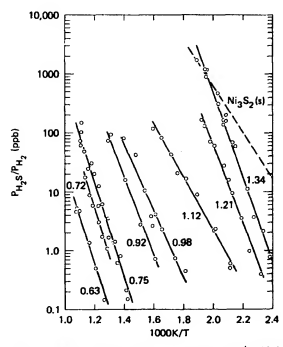


FIG. 3. Sulfur chemisorption isosteres on Ni/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub> with coverage ( $\xi$ ) normalized to CO uptake at 300 K.

TABLE III.  $H_2S/H_2$  isosteres on  $Ni/\alpha-Al_2O_3$  (5 wt%) catalyst (least squares analysis).

Coverage* (ξ)	Temperature range (K)	$\Delta H_f$ (kJ/mol)	ΔS, (J/mol K)	rb	n°
0, 63	863 ± 70	- 159. 4 ± 16.5	-13.5±19.1	0.985	6
0,72	826 ± 50	- 158, 9 ± 11, 5	$-32.0 \pm 13.9$	0.997	6
0.75	806 ± 125	$-155.2 \pm 17.7$	~35.9 ± 22.0	0.994	19
0.92	701 ± 74	- 127. 4 ± 5. 3	-30.0±7.6	0.999	5
0.98	625 ± 85	- 108, 3 ± 25, 8	-13.9 ± 41.3	0.973	9
1, 12	536 ± 75	-86.6±8.3	-7.5 ± 15.5	0.995	10
1.21	478 ± 45	-127.4 ± 12.3	$-117.0 \pm 25.7$	0.995	10
1.35	477 ± 50	$-136.7 \pm 16.0$	$-152.0 \pm 33.5$	0.993	17

\*Coverage normalized to CO uptake (46  $\mu$  mol/g catalyst) at 300 K.

was injected into the recirculating gas, no  $H_2S$  was observed to penetrate the catalyst bed when the coverage was below  $\xi=0.6$ . For successively higher initial coverage, the  $H_2S$  level would quickly rise and then slowly decline with a time constant about 50 min. The decay rate decreased with the decreasing  $H_2S$  level; decay required approximately 4 h at 100 ppb and proportionately longer at lower  $H_2S$  concentration. After injection at an aliquot of  $H_2S$ , the sample was held at the highest experimental temperature for that coverage in order to speed the approach to equilibrium. Slow diffusion through the porous catalyst particles at low  $H_2S$  concentration is most likely responsible for the long times required to reach uniform coverage.

#### IV. DISCUSSION

The results of the present study of sulfur chemisorption thermodynamics for the  $Ni/\alpha$ - $Al_2O_3$  can be described in the following terms. At low coverage ( $\xi < 0.8$ ) chemisorbed sulfur on nickel crystallites yields a heat of formation of -160 kJ mol<sup>-1</sup> and an entropy of 30 J mol<sup>-1</sup> K<sup>-1</sup>. As the coverage increases, the absolute value of the heat of formation decreases because of repulsive interaction

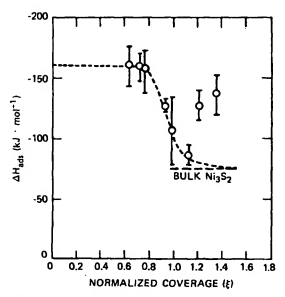


FIG. 4. Heat of adsorption of sulfur via  $H_2S/H_2$  on  $Ni/\alpha-Al_2O_3$  vs coverage ( $\xi$ ) normalized to CO uptake at 300 K.

TABLE IV. H<sub>2</sub>S/H<sub>2</sub> isotherms for Ni/α-Al<sub>2</sub>O<sub>3</sub> (5 wt%) catalyst.<sup>2</sup>

Relative		$P_{ m H_2S}/P_{ m H_2}$ (ppl	o)	Fractional S-saturation
coverage <sup>b</sup>	823 K	623 K	473 K	coverage
0.63	0.46	(4.7×10 <sup>-4</sup> )	•••	0.57
0.72	3.9	$(2.3 \times 10^{-3})$	• • •	0.65
0.75	10,7	(7.4×10°3)	•••	0.68
0.92	(305)	0.78	(3.2×10 <sup>-4</sup> )	0.84
0.98	(715)	4.4	(5.9×10-3)	0.89
1.12	• • •	135	0.67	1.01
1.21	•••	(26, 500)	10.8	
1.35	• • •	(336, 000)	78	

Numbers in parentheses represent extrapolated values. Normalized to CO uptake (46  $\mu$  mol/g catalyst) at 300 K.

between adjacent chemisorbed sulfur atoms. The entropy of the adlayer remains unchanged (within experimental error of  $\pm 12 \text{ J mol}^{-1} \text{ K}^{-1}$ ), until high sulfur coverages are attained (0.63 <  $\xi \le 1.12$ ). At the highest sulfur surface densities encountered in our experiments, a very large entropy change is observed (Table III). An entropy change of this magnitude is well beyond experimental error and is unlikely to be explained by configurational, structural, or vibrational contributions to the entropy of chemisorbed sulfur atoms. The isosteres at  $\xi = 1.21$  and  $\xi = 1.35$  probably represent adsorption of sulfur as SH adspecies or H2S molecules on either the sulfur overlayer on nickel or on the alumina support. These isosteres were measured at low temperatures (below 525 K) where H2S or SH adsorption is more likely to take place. Chemisorption of  $H_2S$  on  $\gamma$  alumina has been reported with isosteric heat of adsorption of 100 to 160 kJ mol $^{-1}$  and  $P_{
m H_{2}8}/P_{
m H_{2}}$  ratios equivalent to 1000 ppb at 525 K.

The energy of the sulfur adsorption bond to nickel surfaces is very large, approaching 465 kJ mol-1, which gives rise to such well known phenomena as sulfur segregation at grain boundaries in the metallurgy of nickel and nickel alloys, and the irreversible poisoning of nickel catalysts by exposure to sulfur containing gases. The heat (enthalpy) of adsorption of sulfur on a nickel surface in equilibrium with a H2S/H2 mixture at a coverage well below saturation is - 158 ± 9 kJ mol-1 at 800 K (with H2S as the reference state for sulfur). The standard thermodynamic reference state for sulfur at this temperature is gaseous diatomic sulfur  $S_2(g)$ , and the heat of formation of gaseous H<sub>2</sub>S at 800 K (Ref. 10) is -89 kJ mol-1. Therefore, the heat of formation of chemisorbed sulfur is -247 ± 9 kJ mol<sup>-1</sup> referenced to diatomic sulfur. The bond strength at 800 K for sulfur adatoms at half saturation coverage is -464 ±9 kJ mol-1, based on -217 kJ mol-1 as the heat of formation of monatomic gaseous sulfur. Bulk Ni<sub>3</sub>S<sub>2</sub>(s) has a heat of formation<sup>10</sup> per mole sulfur of -154 kJ mol<sup>-1</sup> at 800 K; chemisorbed sulfur is therefore 93 kJ mol-1 energetically more stable than the sulfur in Ni<sub>3</sub>S<sub>2</sub>(s), as is readily apparent from Figs. 2 and 3. The heat of formation of sulfur dissolved in nickel [S(b)/Ni] has been measured11 and was found to be - 54 kJ mol-1 over the temperature range from 1173 to 1473 K giving a heat of segregation of -193 kJ mol-1 at approximately half saturation cover-

Correlation coefficient for  $ln(P_{H,S}/P_{H_2})$  vs 1000 K/T.

<sup>&</sup>lt;sup>c</sup>Number of data points in analysis.

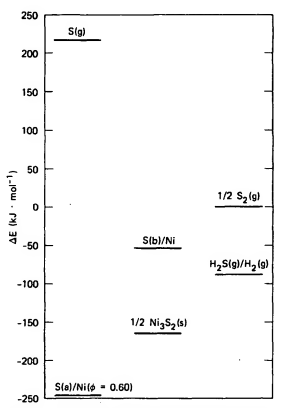


FIG. 5. Energetics of sulfur interaction with nickel at 800 K. S(a): adsorbed sulfur with 0.60 fractional saturation coverage  $(\phi)$ ;  $Ni_3S_2(s)$ : bulk nickel sulfide; S(b): sulfur dissolved in the bulk metal; S(g): gaseous monatomic sulfur. The reference states are  $H_2(g)$ ,  $S_2(g)$ , and Ni(s) at 800 K.

age. Such a large exothermic heat of segregation means that a very low level of sulfur dissolved in the bulk equilibrates with a very high surface coverage. Figure 5 summarizes the energetics of the interaction of sulfur with nickel.

The entropy of adsorption of sulfur on the Ni/α-Al<sub>2</sub>O<sub>3</sub> catalyst surface at about half saturation coverage and 800 K is consistent with the entropy expected for immobile sulfur adatoms. Based on the results of the present study, the entropy change for adsorption is  $-27 \pm 11 \text{ J}$  $mol^{-1}K^{-1}$ , which gives  $56 \pm 11 \text{ J mol}^{-1}K^{-1}$  as the net increase in entropy due to the presence of chemisorbed sulfur on surfaces of the nickel crystallites. Recently, Anderson<sup>12</sup> has measured the frequency of vibration of sulfur adatoms perpendicular to the Ni(100) surface using low energy elastic loss spectroscopy (ELS). ELS is believed 13 to excite only vibrational modes parallel to the incident beam of electrons [in this case normal to the (100) surface]. For this mode, Anderson found a vibrational wave number of 355 cm<sup>-1</sup>. At 800 K, a vibrational mode of this magnitude would contribute 12 J mol-1 K-1 to the entropy of chemisorbed sulfur or 37 J mol-1 K-1 if all three vibrational modes have a frequency of 355 cm<sup>-1</sup>. This leaves only  $19 \pm 11$  J mol<sup>-1</sup> K<sup>-1</sup> to be accounted for by contributions from configuration and other vibrational components. Erley et al. 14 have reported an ELS frequency of 390 cm<sup>-1</sup> for sulfur chemisorption on Ni(111),

a value that gives nearly the same vibrational contribution to chemisorption entropy as determined for Ni(100).

The magnitude of the observed entropy change from adsorption of sulfur on nickel surfaces in equilibrium with gaseous  $\rm H_2S/H_2$  gas mixtures rules out  $\rm H_2S$  and SH as the adsorbed species at moderate coverage. At 800 K, we would expect to observe roughly -180 and -100 J mol<sup>-1</sup> K<sup>-1</sup> for  $\Delta S$  of chemisorbed  $\rm H_2S$  and HS species, respectively. These values are clearly beyond the experimental uncertainty. However, at 500 K, we would expect  $\Delta S = -190$  and -120 J mol<sup>-1</sup> K<sup>-1</sup>, respectively, for chemisorbed  $\rm H_2S$  and HS. We note that the entropy change for the SH species is closer to the observed value,  $-135 \pm 25$  J mol<sup>-1</sup> K<sup>-1</sup>, at high coverage (Table III).

The results of our study of the thermodynamics of sulfur chemisorption on powdered nickel, y-Al<sub>2</sub>O<sub>3</sub>-supported nickel, and  $\alpha - Al_2O_3$ -supported nickel show that sulfur is an even more tenaciously bound catalyst poison for nickel than previously suspected. At 575 K, a typical temperature for catalytic methanation of synthesis gas. our data gives 0.7 ppb H2S in 1 atm H2 as the equilibrium sulfur activity at 90% saturation coverage. Thus, to prevent sulfur accumulation on the catalyst the upper limit of H<sub>2</sub>S in syn gas for methanation is extremely low. At the same time regeneration of a sulfur-poisoned catalyst by hydrogen reduction is impractical because of the low equilibrium partial pressure of H2S. A prohibitive volume of H<sub>2</sub> would be required at temperatures below 700 K to remove chemisorbed sulfur from nickel surfaces. At higher temperatures Ni crystallite sintering would become a problem.

In order to compare our isosteric results with previous measurements,  $^{2,3,5}$  we need to establish reference points for the surface coverage of sulfur on different Ni surfaces and topographies. Perdereau and Oudar<sup>2</sup> reported that the saturation sulfur coverage on nickel surfaces (800 ppm  $\rm H_2S$  in  $\rm H_2$  at 875 K after 2700 s exposure) remains nearly constant (8.2 ± 0.3×10<sup>14</sup> atoms per cm<sup>2</sup>), regardless of orientation of the crystal plane. For Ni(100) single crystal planes the saturation coverage is  $8.0 \times 10^{14}$  atoms per cm<sup>2</sup>, which represents 50% of the density of nickel atoms at the surface (15.9×10<sup>14</sup> atoms per cm<sup>2</sup>). This result is consistent with the C(2×2) LEED pattern observed<sup>2,15</sup> for saturation with sulfur on

TABLE V. Saturation coverage for sulfur chemisorption on nickel.

Material	T (K)	$P_{\mathrm{H}_2\mathrm{S}}/P_{\mathrm{H}_2}$ (ppm)	S/Hª	Ref.	Equivalent <sup>b</sup> § - S/CO
Ni foil	1173	100		2	1.02
Ni foil	1073	150	• • •	2	1.12
Ni foil	973	150	• • •	2	1. 13
Ni/MgO · Al <sub>2</sub> O <sub>3</sub>	918	8	0.74	3	1,00
Ni/MgO · Al <sub>2</sub> O <sub>3</sub>	823	2	U. 74	3	1.05
Ni powder	723	25	1.14	5	1, 15
3 wt2 Ni/Al <sub>2</sub> O <sub>3</sub>	723	25	1.04	5	1.15
16 wt'& Ni+0.5 wt'& Pt/Al2O3	723	25	0.82	5	1. 15
16 wt's Ni-0.5 wt's Pt/Al2O3	793	25	0.74	ā	1.14

<sup>a</sup>Saturation S coverage vs H<sub>2</sub> chemisorption from reference. <sup>b</sup>¢ equivalent S coverage vs CO chemisorption, this work.

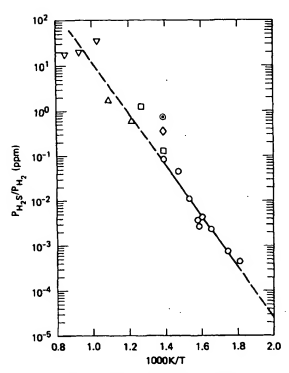


FIG. 6. Sulfur activity vs reciprocal temperature for chemisorbed sulfur. (v) Ni foil (Ref. 2); ( $\Delta$ ) Ni/MgO<sub>3</sub>·Al<sub>2</sub>O (Ref. 3); ( $\Phi$ ) Ni powder (Ref. 5); ( $\Phi$ ) Ni/Al<sub>2</sub>O<sub>3</sub> (Ref. 5); ( $\Phi$ ) Ni/ $\Phi$ -Al<sub>2</sub>O<sub>3</sub> (this work); ( $\Phi$ ) least squares analysis (this work). All points are for fractional saturation coverage ( $\Phi$ ) = 0.88.

the Ni(100) plane. A number of other workers<sup>16-20</sup> report similar values for  $H_2S$  adsorption at 200 to 300 K on nickel films and supported nickel.

However, in the determination of the surface site density on Ni surfaces by means of  $H_2$  or CO adsorption, the adsorption stoichiometries are not firmly established beyond a factor of 2. Rostrup-Nielsen, <sup>3</sup> using a H/Ni(s) = 0.73, arrived at a value of S/Ni(s) = 0.54 at saturation coverage on Ni/MgO·Al<sub>2</sub>O<sub>3</sub>. Based on an average surface site density of 15.4×10<sup>14</sup> Ni atom/cm<sup>2</sup> for the low index planes of Ni, the sulfur saturation density (8.3 ×10<sup>14</sup> S atom/cm<sup>2</sup>) so calculated is in good agreement with Perdereau's results (8.2×10<sup>14</sup> S atom/cm<sup>2</sup>). By contrast the results of Pannell *et al.*, <sup>21</sup> in which H/Ni(s) is assumed to be unity, yields sulfur saturation coverage corresponding to 11.4×10<sup>14</sup> S atom/cm<sup>2</sup> for powdered Ni and 17.6×10<sup>14</sup> S atom/cm<sup>2</sup> for the alumina-supported Pt-promoted Ni catalyst (16 wt% Ni, 1.5 wt% Pt).

In our work CO adsorption was employed as a measure of Ni surface site density. On well defined Ni planes CO adsorption studies  $^{22-26}$  gave nearly constant saturation coverage, corresponding to  $11.0\pm0.3\times10^{14}$  CO molecules/cm² regardless of crystal orientation. Based on CO adsorption we arrive at a sulfur saturation coverage corresponding to  $12.2\pm0.8\times10^{14}$  S atoms/cm² on 5 wt% Ni/ $\alpha$  -Al<sub>2</sub>O<sub>3</sub>. On the basis of the resulting adsorption ratio S/CO=1.11 we have evaluated the fractional sulfur saturation coverage ( $\phi$ ) for this catalyst (Table IV), and the equivalent saturation coverage corresponding to saturation conditions used in previous work (Table V).

Now we are in a position to compare the results reported by various authors with our S chemisorption isosteres. In Fig. 6 we have plotted the data near saturation coverage ( $\phi=0.88$ ). It will be noted that over a wide range of temperatures the various data points are in satisfactory agreement with our results represented by the solid line. However, a similar comparison at lower S coverage ( $\phi=0.7$ ) demonstrate that our equilibrium  $\rm H_2S/H_2$  partial pressure ratios are considerably lower than those reported previously. <sup>2.3</sup> Based on our observation that the approach to equilibrium is very slow at low  $\rm H_2S$  concentration, we suggest that equilibrium may not have been reached in the earlier studies.

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## The poisoning effect of sulfur-containing impurity gas on a SOFC anode: Part I. Dependence on temperature, time, and impurity concentration

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#### Abstract

The poisoning effect of a sulfur-containing impurity on the electrochemical oxidation of  $H_2$  has been studied at 1023-1273 K in a gas mixture of  $H_2$  and  $H_2O$  at the interface between a Ni-YSZ cermet electrode and a YSZ electrolyte using a complex impedance analysis and a DC polarization method. The polarization resistance and the overvoltage of the electrode increased when the  $H_2S$  concentration exceeded 0.05, 0.5, and 2 ppm at 1023, 1173 and 1273 K, respectively. A large temperature dependence was observed, which insists that a high-grade desulfurization is necessary at lower operating temperatures. The time needed for the influence of the sulfide impurity to saturate was almost independent of the sulfide concentration, and was found to be approximately 12, 9 and 4 ks at 1023, 1173 and 1273 K, respectively. Within the present experimental conditions, the performance loss caused by the sulfur-poisoning was recoverable when the sulfur source was removed from the fuel; the time needed for the recovery was approximately 360, 90 and 4 ks at 1023, 1173 and 1273 K, respectively. The degree of sulfur-poisoning was found to depend not on the equilibrium partial pressure of  $S_2$  but on the total sulfur content in the fuel. © 2000 Elsevier Science B.V. All rights reserved.

Keywords: Solid oxide fuel cell; Fuel electrode; Sulfur; Poisoning; Impurity; Complex impedance

#### 1. Introduction

Fuel cells have high fuel-to-electricity conversion efficiencies and low levels of NO<sub>x</sub> and SO<sub>x</sub> emission. Solid oxide fuel cells (SOFCs) have attracted much attention because of their distinguished advantages over other types of fuel cells such as high electric efficiency, high-quality exhaust heat and system

Among several types of SOFCs, electrolyte-supported cells using yttria-stabilized zirconia (YSZ) as the electrolyte have many material problems remain-

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compactness. High operating temperatures of SOFCs ranging from 873 to 1273 K enables them to use a wide variety of fuels including natural gas. For natural gas fueled SOFCs, however, the effect of sulfide impurities must be taken into account and desulfurization was considered since several parts per million (ppm) of sulfur-containing impurities are usually added as odorant to natural gas distributed to customers by pipelines.

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ing to be solved which arise from the high operating temperature, typically around 1273 K. A reduction in the operating temperature leads to the benefits which include low-cost metallic separators instead of more expensive ceramic ones [1], better long-term performance stability, and system compactness by reducing the thickness of the heat insulator. Fundamental research, therefore, has been carried out on manufacturing and testing the cells reduced-temperature operation using electrode-supported cells [2–5] with a thin YSZ electrolyte film on thick porous electrodes as well as the electrolyte-supported cells in which the high oxide-ion conductor of doped LaGaO<sub>3</sub> is used as the electrolyte [6–8].

Although many benefits are expected by reducing the operating temperature, it should be considered that at reduced temperatures sulfide impurities may easily adsorb on the surface of a SOFC fuel electrode to degrade the resistance of the electrode against the sulfide impurities. Of impurities in fuels, the sulfurcontaining compounds, which are present primarily as hydrogen sulfide (H<sub>2</sub>S) under the fuel electrode environment are expected to have the greatest impact on SOFC performance [9].

In previous studies on the poisoning effect of sulfur, a direct-current polarization method and complex impedance analysis were used to evaluate cell performance loss, which showed that even low levels in ppm of H<sub>2</sub>S may cause significant performance losses at the fuel electrode of SOFC. Geyer et al. [10] reported that the polarization resistance of a cermet electrode increased by a factor of two by addition of only 5 ppm of H<sub>2</sub>S at 1223 K in H<sub>2</sub> with 3% H<sub>2</sub>O. If the H<sub>2</sub>S content is low, however, the performance recovers upon removing the sulfur impurity from the fuel [11-13]. As for the high levels of H<sub>2</sub>S concentration, Dees et al. [14] reported that the polarization resistance increased by a factor of two by addition of 105 ppm H<sub>2</sub>S at 1273 K in H<sub>2</sub> with 3% H<sub>2</sub>O and the poisoning effect was irreversible. Several possible mechanisms may account for the decrease in the performance of the electrode due to the presence of H<sub>2</sub>S in the fuel. The exact mechanism responsible for the performance loss, however, is not clearly defined because of the lack of detailed experimental data. In this study, we have investigated the time-constant of sulfur saturation and removal, and the dependence of the poisoning

effect on the operating temperature, the sulfur contaminant concentration and the equilibrium partial pressure of  $S_2$  produced in fuels containing  $H_2S$ .

#### 2. Experimental

#### 2.1. Test cell preparation

Dense pellets (20 mm in diameter and 2 mm in thickness) of 8 mol% YSZ were used as the electrolytes for electrochemical cells. NiO powder with an average grain size of 0.9 µm (Nihon-Kagaku-Sangyo) was dispersed in a mixed toluene solution of zirconium octylate (Zr(C<sub>4</sub>H<sub>9</sub>CH (C<sub>2</sub>H<sub>5</sub>)CO<sub>2</sub>)<sub>4</sub>, 99% pure, Nihon-Kagaku-Sangyo) and yttrium octylate  $(Y(C_4H_9CH(C_2H_5)CO_2)_3,$ 99% pure, Nihon-Kagaku-Sangyo), the mixing ratio of which was adjusted to the ratio corresponding to the composition of 8 mol% Y<sub>2</sub>O<sub>3</sub>-92 mol% ZrO<sub>2</sub>. The paste thus prepared was screen-printed onto one side of the electrolyte pellet, followed by firing at 1723 K for 7.2 ks to obtain a porous NiO-YSZ composite [15]. The thickness and porosity of a Ni-YSZ cermet electrode obtained after reduction of the NiO-YSZ composite were about 25 µm and 45%, respectively. The YSZ content in the cermet electrode is 10 wt%. For a counter electrode, Pt paste (Tokuriki Kagaku 8103) was painted symmetrically to the Ni-YSZ cermet electrode on the other side of the pellet and fired at 1273 K for 7.2 ks. For a reference electrode, the Pt paste was painted at the perimeter of the pellet and fired at 1273 K for 7.2 ks.

#### 2.2. Electrochemical measurement

Fig. 1 shows the test geometry used in the electrochemical measurements. Two Sr-doped LaCrO<sub>3</sub> plates with grooves were used as both manifolds and current collectors. Humidified H<sub>2</sub> fuel was fed into the doped LaCrO<sub>3</sub> plates from the top of a furnace and was supplied to the reference, counter and the Ni-YSZ cermet electrode. H<sub>2</sub>S was added to the fuel as the sulfide impurity at the concentration range from 0.02 to 15 ppm. Platinum wires were used as voltage terminals for the working and reference electrodes.

The complex impedance measurements were made

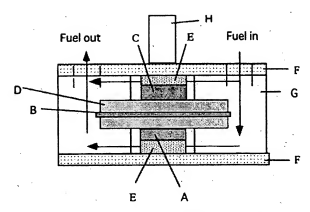


Fig. 1. Test geometry used in the electrochemical measurements. A: working electrode, B: reference electrode (Pt), C: counter electrode (Pt), D: YSZ pellet, E: Pt mesh for current collection, F: doped LaCrO<sub>3</sub> plate for current collection and manifold, G: Al<sub>2</sub>O<sub>3</sub> plate for manifold, H: pushrod.

over the frequency range of 1 mHz-100 kHz with an applied amplitude of 10 mV using the three-terminal method at an equilibrium potential. The system for the impedance measurements was composed of a frequency response analyzer (Solartron 1260), a potentiostat (Solartron 1287) and a personal computer (Compac Armada I 120T). In order to keep the electrode stable during the complex impedance measurements, the electrode was pretreated under a DC polarization at the current density of 0.3 A cm<sup>-2</sup> in the fuel without H<sub>2</sub>S for 180 ks followed by annealing for 180 ks at the equilibrium potential at each temperature. The steady-state polarization was measured by a current interruption method using a current-pulse generator (Hokuto Denko HC-110). Also before the polarization measurements, pretreatment was conducted at current density of 0.3 A cm<sup>-2</sup> in the fuel without H<sub>2</sub>S for 180 ks.

#### 3. Results and discussion

#### 3.1. Poisoning by sulfur and recovering at 1273 K

Fig. 2 shows the change in the complex impedance spectrum with time when  $H_2S$  is added to the fuel, 79%  $H_2 + 21\%$   $H_2O$ , at 1273 K. The  $H_2S$  concentration,  $C(H_2S)$ , was set at 1, 2, 4, 6, 8, 10 and 15 ppm, and the impedance measurements were

repeated consecutively until the spectra showed no change with time. The figure shows the four selected time-evolutions of the impedance spectra. Two arcs seemed to be overlapped in the impedance plots which show depressed semi-circles. As shown in these figures, the addition of H<sub>2</sub>S did not change the size of impedance arcs at the concentration of 1 ppm, but increased the size of impedance arcs at 2 ppm and more. These results indicate that the Ni-YSZ cermet electrode was degraded by the sulfide impurity when the value of C(H<sub>2</sub>S) exceeded 2 ppm at 1273 K. It is found that the size of impedance arcs initially increased with time at C(H<sub>2</sub>S) of 2 ppm or more and saturated at time between 3.6 and 4.8 ks after the H<sub>2</sub>S addition. Fig. 3 shows the complex impedance spectra after full equilibration with the fuels with several concentrations of H<sub>2</sub>S. The steadystate impedance curves became larger with increase of C(H<sub>2</sub>S), but no significant difference appeared in the shape of the curves even when C(H<sub>2</sub>S) increased. Fig. 4 shows the change of the complex impedance spectra after the fuel with 15 ppm H<sub>2</sub>S was switched to a H<sub>2</sub>S-free fuel. The time in the legend is the time elapsed after H<sub>2</sub>S was removed from the fuel. It is found that the size of the impedance arcs decreased with time and recovered to its original level at about 3.6 ks after the H<sub>2</sub>S removal. The complex impedance spectra of the electrode equilibrated in the fuel with 2-10 ppm H<sub>2</sub>S were also found to recover to the original size and shape at about 3.6 ks after the H<sub>2</sub>S was removal. From these results it is concluded that 2-15 ppm H<sub>2</sub>S impurity deteriorates the electrode at 1273 K, but this poisoning effect is reversible if the H<sub>2</sub>S-free fuel is supplied again.

Fig. 5 shows the overvoltage (with IR) of the electrode as a function of time at the current density of 0.3 A cm<sup>-2</sup> (a) and (b) show the overvoltage after H<sub>2</sub>S was added to the fuel and H<sub>2</sub>S was removed from the fuel, respectively. The overvoltage started to increase with time when H<sub>2</sub>S was added to the fuel and saturated after a sufficient time; the steady-state overvoltage was larger than that in the H<sub>2</sub>S-free fuel by about 2 mV at C(H<sub>2</sub>S) of 2 ppm and 74 mV at C(H<sub>2</sub>S) of 15 ppm. The degraded electrode performance fully recovered upon switching to the H<sub>2</sub>S-free fuel. These results are consistent with the results in the polarization resistance evaluated by the complex impedance analysis.

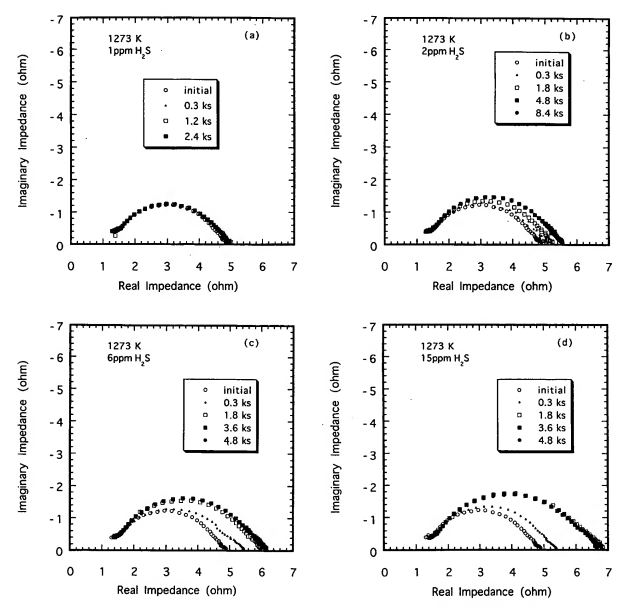


Fig. 2. The frequency dispersion diagram of the complex impedance spectra of the electrode in the fuel with  $H_2S$  of (a) 1, (b) 2, (c) 6 and (d) 15 ppm at 1273 K.

#### 3.2. Temperature dependence of sulfur poisoning

Fig. 6 (a) and (b) show the complex impedance spectra after equilibration in the fuels with different  $H_2S$  concentrations at 1173 and 1023 K, respectively. At 1173 K, the impedance arcs increased when the  $C(H_2S)$  exceeded 0.5 ppm, which was lower than

that 2 ppm at 1273 K. At 1073 K the impedance arcs began to increase at a much lower concentration of 0.05 ppm. These results suggest that reducing the operating temperature degrades the resistance of the electrode against the sulfide impurities to cause the poisoning at lower sulfur concentrations.

Fig. 7 shows typical impedance spectra measured

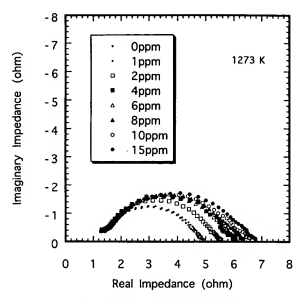


Fig. 3. The complex impedance spectra measured after equilibration in the fuels with several concentrations of H<sub>2</sub>S at 1273 K.

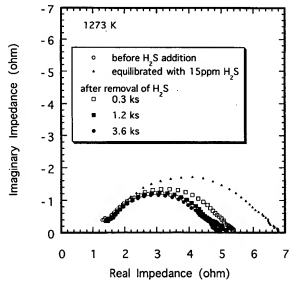


Fig. 4. The change of the complex impedance spectra at 1273 K when 15 ppm  $H_2S$  was removed from the fuel.

at 1173 K; (a) and (b) show the change of impedance spectra after addition and removal of 1 ppm  $H_2S$ , respectively. When the  $H_2S$  was added to the fuel, the size of the impedance arcs increased with time and saturated after approximately 9 ks which was longer than at 1273 K. When  $H_2S$  was removed from

the fuel, the size of the impedance arcs decreased with time and finally recovered to its original level before the poisoning. However, a longer time, approximately 90 ks was needed for the recovery at 1173 K as compared with that at 1273 K, which became more remarkable at a lower temperature. The typical impedance spectra measured at 1023 K are shown in Fig. 8(a) and (b). At 1023 K, it took approximately 12 ks for the size of the impedance arcs to saturate after the H<sub>2</sub>S addition and took approximately 360 ks for the recovery after removing H<sub>2</sub>S. It is also found that the recovered impedance arc is smaller than that before H<sub>2</sub>S addition, the cause of which has not yet been identified.

Fig. 9 shows the difference between the steady-state overvoltage in the fuel with the  $H_2S$  impurity and the initial overvoltage without  $H_2S$  impurity as a function of  $C(H_2S)$  and the operating temperature at the current density of 0.3 A cm<sup>-2</sup>. The minimum concentrations of  $H_2S$  which caused the increase of the overvoltage showed good correspondence to those for the polarization resistance evaluated by the complex impedance analysis, i.e. 2, 0.5 and 0.05 ppm at 1273, 1173 and 1023 K, respectively. Similar to the results in the complex impedance measurements, the increase in the overvoltage recovered at all temperatures when  $C(H_2S)$  returns to 0 ppm, i.e. the poisoning effect of sulfur on the overvoltage is also reversible over the concentration range studied.

The degree of the poisoning caused by sulfur was thus found to have a large dependence on the operating temperature; the critical sulfur level for the performance loss is lower at lower operating temperatures, and at a selected sulfur concentration, the overvoltage increases more largely at lower temperatures. Since it can be presumed that the sulfur impurity adsorbs more easily on the electrode as temperature decreases, these results suggest that an adsorbed impurity inhibits some processes in electrochemical reactions at the interface between the fuel electrode and the electrolyte.

#### 3.3. Dependence on $S_2$ partial pressure

The partial pressure of  $S_2$ ,  $P(S_2)$ , in the fuel is determined by the following equilibrium reactions as a function of  $H_2$  partial pressure,  $P(H_2)$ ,  $C(H_2S)$  and temperature.

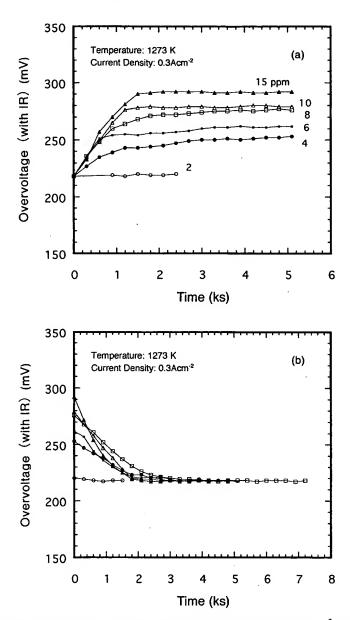


Fig. 5. The overvoltage (with IR) of the electrode as a function of time at a current density of 0.3 A cm<sup>-2</sup> after (a) addition and (b) removal of  $H_2S$ .

$$2H_2S = 2H_2 + S_2$$
 (1)  $2H_2O = 2H_2 + O_2$  (2)

Since  $P(H_2)$  in the fuel is determined by the following equilibrium reaction as a function of the partial pressure of  $H_2O$ ,  $P(H_2O)$ , and the partial pressure of oxygen,  $P(S_2)$  is also a function of these partial pressures.

To discuss the influence of  $P(S_2)$  on the sulfur poisoning of the electrode, we measured the steady-state polarization as a function of  $P(S_2)$  which was controlled by changing  $P(H_2O)$  with constant  $C(H_2S)$  or by changing  $C(H_2S)$  with constant

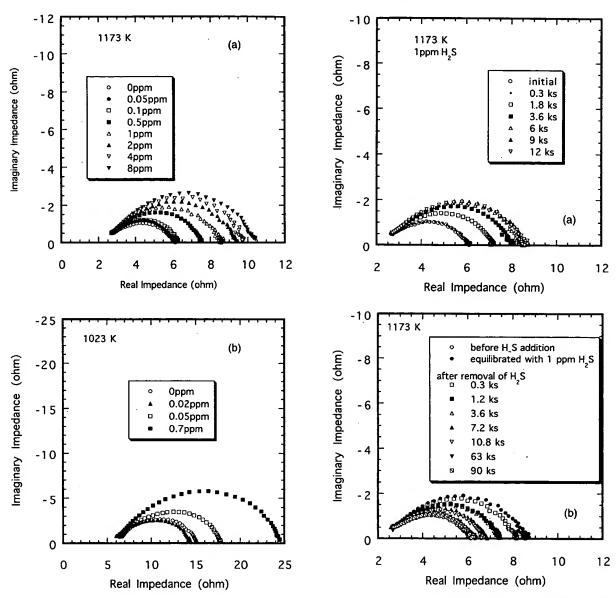
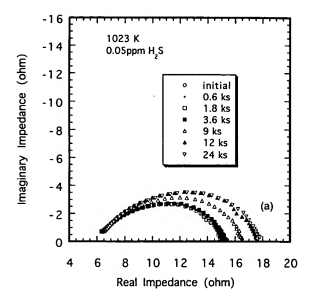


Fig. 6. The complex impedance spectra measured after equilibration in the fuels with several concentrations of H<sub>2</sub>S at (a) 1173 K and (b) 1023 K.

Fig. 7. Typical complex impedance spectra measured at 1173 K after (a) addition and (b) removal of 1 ppm  $H_2S$ .

 $P(\rm H_2O)$ . Fig. 10 shows the increase of the overvoltage,  $\Delta V$ , by the poisoning as a function of  $P(\rm S_2)$  at the current density of 0.3 A cm<sup>-2</sup>. The closed circles indicate plots with constant  $P(\rm H_2O)$  with the value of 21% at all temperatures. With the constant  $P(\rm H_2O)$ ,  $\Delta V$  increased with  $P(\rm S_2)$ , which can be regarded as caused by the increase of  $P(\rm S_2)$  or

 $C(\mathrm{H_2S})$ . There is no previous study in which the effect of  $C(\mathrm{H_2S})$  on fuel electrodes is separated from that of  $P(\mathrm{S_2})$  to identify the dominant factor of sulfur-poisoning.  $\Delta V$  was, therefore, measured with constant  $C(\mathrm{H_2S})$  in order to separate the influence of  $C(\mathrm{H_2S})$ . The open circles indicate plots with constant  $C(\mathrm{H_2S})$ . The values of  $C(\mathrm{H_2S})$  in the constant  $C(\mathrm{H_2S})$  conditions at 1023, 1173 and 1273 K were



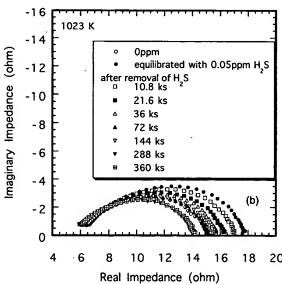


Fig. 8. The typical impedance spectra measured at 1023 K after (a) addition and (b) removal of 0.05 ppm  $H_2S$ .

0.05, 0.5 and 4 ppm, respectively. It can be seen that the  $P(S_2)$  dependence of  $\Delta V$  in the fuel with constant  $C(H_2S)$  is different from that in the fuel with constant  $P(H_2O)$  at all temperatures. With the constant  $C(H_2S)$ ,  $\Delta V$  showed little dependence on  $P(S_2)$ . These results indicate that a factor dominating the poisoning by the sulfide impurity is not  $P(S_2)$  but the total sulfur content in the fuel. In order to obtain

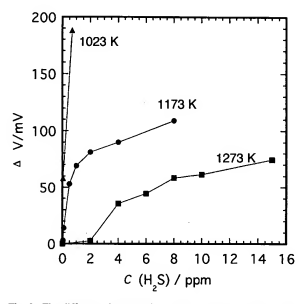


Fig. 9. The difference between the steady-state overvoltage in the fuel with  $\rm H_2S$  and the initial overvoltage in the  $\rm H_2S$ -free fuel as a function of  $\rm \it C(H_2S)$  at the current density of 0.3 A cm<sup>-2</sup>.

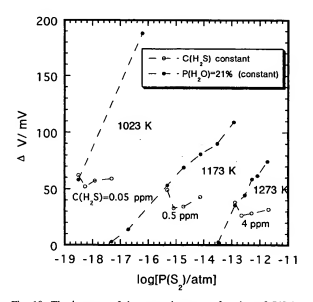


Fig. 10. The increase of the overvoltage as a function of  $P(S_2)$ ; the open and closed circles indicate plots with constant  $C(H_2S)$  and with constant  $P(H_2O)$ , respectively. The values of the constant  $C(H_2S)$  are 0.05, 0.5 and 4 ppm for 1023, 1173 and 1273 K, respectively, and the value of the constant  $P(H_2O)$  is 21% at all temperatures.

further information for investigation of the mechanism of the poisoning, an equivalent circuit analysis of the impedance spectra has been conducted [16].

#### 4. Conclusions

The influence of the sulfide impurity on the electrochemical properties of the fuel electrode was investigated by using the complex impedance analysis and the DC polarization method. The polarization impedance and the DC overvoltage of the electrode increased when H<sub>2</sub>S concentration exceeded 0.05, 0.5, and 2 ppm at 1023, 1173 and 1273 K, respectively. The time needed for the influence of the sulfide impurity to saturate was almost independent of the sulfide concentration, and was found to be about 12, 9 and 4 ks at 1023, 1173 and 1273 K, respectively. Within the experimental conditions, the performance loss caused by the sulfur-poisoning was recoverable when H<sub>2</sub>S was removed from the fuel; the time needed for the recovery was found to be 360, 90 and 4 ks at 1023, 1173 and 1273 K, respectively. Thus, the poisoning effect has been found to have a large dependence on the operating temperature, which insists that a high-grade desulfurization will be necessary for reducing the operating temperature. From the dependence of sulfur poisoning on the equilibrium partial pressure of S<sub>2</sub> we have reached the conclusion that the sulfide poisoning was governed by the total sulfur content in the fuel.

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# Activity of Nickel Catalysts for Steam Reforming of Hydrocarbons

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by the composition of the catalyst. The specific activity is very low on some support observed for reforming of ethane is generally following that obtained for reforming A number of catalysts have been investigated to identify some factors of chemical materials, and when alkali is present. The activity correlates with the surface heterogeneity expressed by adsorption of nitrogen on the nickel surface. The activity trend composition and surface structure which may influence the activity for steam reforming of hydrocarbons. The kinetics for ethane reforming at 500°C are influenced of other hydrocarbons, hydrogenolysis of ethane, and methanation of carbon monoxpresence of alkali. Various explanations for the effect of the carrier and of alkali ide; whereas decomposition of ammonia appears unaffected by the carrier

many kinetic studies of chemical or a given conversion in industrial reactors. ionversion. Steam reforming of higher hydiocarbons is a complex process including ocesses a single catalyst sample is invesgated in great detail to provide a basis for culation of the catalyst volume required ubular steam reforming of higher hydroeveral consecutive and parallel reactions sence comparable to that on the overall arbons is a process where secondary eficts of catalyst activity may have an iniverall reaction can be described by which some may result in coke. reakdown of the hydrocarbon:

Ξ followed by establishment of the equilibria:  $C_nH_m + nH_2O \rightarrow nCO + \left(n + \frac{m}{2}\right)H_2$ 

ଚ  $CO + H_2O = CO_1 + H_2$ 

3  $CO + 3H_2 = CH_4 + H_2O.$  Tubular reformers are working normally at talyst inlet and exit temperatures in the finimum exit temperature to be applied without breakthrough of higher hydrocar-450-550 and 650-850°C, respeclively. The catalyst activity determines the ranges

reformers for production of synthesis gas reactions (2) and (3) established at the bons. This may be a problem in reformers for towns gas production (I, 2). In most and hydrogen, the composition of the exit gas will be very close to the equilibria of exit temperature and, normally, design restrictions to ensure long tube lives will determine the maximum throughput.

smaller (1, 3). However, this study deals to evaluate a complete kinetic expression to be applied in reactor design. As shown High catalyst activity may result in cause the breakdown of the higher hydrocarbons may be completed in the colder part of the bed where coking rates are only with the activity for conversion of the higher hydrocarbons, reaction (1), whereas considered and no attempts have been made it may imply reduced risks of coking bethe selectivity problems will be reported ous reactions, (2) and (3), have not been in Table I very different results have been andin (4) observed the reaction order with lower tube wall temperatures. In addition later. Moreover, the rates of the simultane-(4-8). Balashova, Slovokhotova and Balreported on the kinetics of reaction (1

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TABLE 1
RESULTS FROM SOME KINETIC STUDIES OF STEAM REFORMING OF HIGHER HYDROCARBONE

	Catalvet		Tom	D	Kii	Kinetic coefficients	Activati
Authors	system	Hydrocarbon	(C)	(atm abs) $\alpha_{C_nH_m}$	αC <sub>n</sub> Bm	αH <sub>2</sub> 0	energy (kcal/m
Balashova, Slovok- hotova and	Ni/SiO <sub>2</sub> · Ni/C	Cyclohexane 400-460	400-460	(1)	O N g o	0 0-1 No activity	22-24
Balandin (4) Bhatta and Dixon (6) $Ni/\gamma$ -Al <sub>2</sub> O <sub>3</sub>	Ni/y-Al,O3	n-Butane	425-475	30	0	, <del>-</del>	C E I
Bhatta and Dixon (6)	$Ni/\gamma-Al_2O_3$	n-Butane		30	0	٠,	13%
	$Ni/\alpha$ - $Al_2O_3UO_2$		404-491	30	_	9.0-	24.
Phillips, Mulhall and Turner (?)	Ni/y-Al <sub>2</sub> O <sub>3</sub>	n-Hexane	360-450	15	0.3	0-0.04	213
Saito et al. (8)	Ni/SiO <sub>2</sub>	n-neptane n-Butane	370-450		0	-	1

respect to steam to vary with the range of steam partial pressure, and the data of Bhatta and Dixon (6) demonstrated that the kinetics may change significantly from catalyst to catalyst.

The purpose of the present work has been to identify some of the factors in chemical composition and surface structure of the catalyst which may influence the activity. Various catalyst preparations have been investigated but a more detailed kinetic study has only been made with a single catalyst sample. The study has been limited to temperatures around 500°C, a typical inlet temperature of the catalyst bed, because a high activity for reaction (1) is assumed to be important in this part of the bed. The activity has been estimated mainly from isothermal steam reforming

experiments at atmospheric pressure. The ane, being the simplest higher hydrocarbon, was selected as feed. Some experiments have been performed at pressure using various pure hydrocarbons and naphthal feed. Finally, the activities for other reactions, methane reforming, methanation of composition of ammonia and ethane hydrocarposis, have been estimated for some of the catalysts from experiments at atmospheric pressure. The activity tests have been supplemented by chemisorption studies.

### METHODS

# 1. Ethane Reforming at Atmospheric Pressure

Apparatus. The experiments were performed in a simple flow system with a

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Fig 1. Apparatus for studies of ethane reforming and other reactions at atmospheric pressure. 👬

changed during the run was varied unsyste-

ubular fixed bed reactor as shown in Fig. 1.
The reactor, which was made of 18/8 stainless steel, was surrounded by an electrically leasted furnace. A metal block with an excitant diameter of 25 mm was mounted on the reactor to ensure isothermal conditions. The internal diameter of the reactor was 5.04 mm, and a thermowell with external diameter of 2.0 mm was placed in the axis of the reactor. The temperature of the catalyst bed was adjusted by means of a high ristor controller. The axial temperature diameter is estimated to be less than 1°C/

fin the catalyst bed.
The gas flows were controlled by flowelers, whereas the addition of H<sub>2</sub>O was

placed in the reactor. The catalyst was ormally, the activity stabilized within less gan 30 min. However, over periods of days sed for two runs. The parameter to be füced catalyst as 0.3-0.5 mm particles was heated in H2 (0.18 mol/hr) to the temperature to be used for the experiment  $H_2O/H_2 = approx 10$ ) to avoid oxidation of the catalyst. When the exit gas flow was stabilized a sample of the dry exit gas was ould be ascribed to sulfur poisoning. Owing pprox 6 hr. In a few cases the bed was Gases. All gases except H2 were taken ess than 0.03%) was made in an Oerlikon ation. C2He was purified over NiMo graphic analyses showed impurities of CH, After condensate was observed in the cooler and steam flow checked by sampling of ments, also CO2 were added. All experinents were performed with H2 in the feed gradual deactivation was observed which prinally having a maximum duration of estalyst and ZnO at 270°C. Gas chromato-Hs (0.02-0.08), and C3Hs (0.02-0.40 vol condensate, ethane and, in a few experitaken, and the conditions were changed. this the bed was changed after each run, . N<sub>2</sub> was purified over Cu wire at 250-Procedure. A sample of 0.1-0.4 g of rem cylinders. H<sub>2</sub> (impurities (mainly O<sub>2</sub>) lectrolyzer and used without further purimax 0.01 vol %), C2H4 (0.04-0.66 vol %) 275°C. CO<sub>2</sub> was used without purification. \*(normally 500°C) and steam was added lowed on a buret.

matically with time to minimize the influence of poisoning. Moreover, as a check each experiment was completed at the same conditions as the starting conditions. After each experiment the weight of the catalyst was checked.

The gas sample was analyzed by means of Orsat (CO<sub>2</sub>) and gas chromatographic analysis. At the normal conditions the main products were CO<sub>2</sub> and H<sub>2</sub>.

Evaluation. The reaction of ethane with steam:

$$C_2H_6 + 4H_2O = 2CO_2 + 7H_2,$$
 (4)

can be considered irreversible as K (500°C) = 155 atm<sup>4</sup>.

For simplicity a kinetic expression of the power form was selected:

$$r = F \frac{dx}{dw} = A e^{-B_o l RT} \prod_i p_i \alpha_i, \quad (5)$$

where  $F = C_2H_6$  inlet flow, W = catalyst weight, and x = conversion. The conversion of  $C_2H_6$  was calculated on basis of the exit gas analysis by:

$$x = \frac{(\text{CH}_4) + (\text{CO}) + (\text{CO}_2)}{(\text{CH}_4) + (\text{CO}) + (\text{CO}_2) + 2(\text{C}_2\text{H}_6)} \times 100\%. (6)$$

Relative deviations in ethane The kinetic parameters were estimated from temperature gradients. On the basis of the eters were changed until the variance was minimized. In the calculations the meaculated on the basis of the exit gas analysis This procedure was found to be more accurate than using the readings of the flowmeters. The mass balance of the apparatus was checked in a few experiments where gas samples were taken before and after Blank experiments at 500°C with alumina in question, and the estimated values were suming no diffusion restrictions and no calculated rate constants the kinetic paramsured  $H_2O$  addition was used as the  $H_2O$ ieed, whereas the inlet gas flows were caland the measurement of the exit gas flow. plots of the conversion and the parameter used for computer integrations of (5) ascontents of less than 1% were indicated. particles showed no conversion. the reactor.

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analysis and the exit flow. The conversion C2H6 and CO2 in the feed, and in all these and after the reactor. The inlet gas flows of CO2 and C2H6 were calculated from the CO<sub>2</sub>/C<sub>2</sub>H<sub>6</sub> ratio of the feedgas, the exit gas procedure described could not be applied in experiments with experiments gas samples were taken before was calculated from the ethane mass calculation balance.

the initial rate ri was calculated at followcatalysts with different kinetic parameters In order to compare the activities of standard conditions: H2O/C2H6 =  $H_2O/H_2 = 10$ ; temperature = 500°C; and pressure = 1.0 atm abs.

## 2. Other Reactions Tested at Atmospheric Pressure

sition, and ethane hydrogenolysis. CH4 and NH3 were taken from cylinders. CH4 concarbons was purified by washing in a The apparatus used for ethane reforming experiments was also used for experiments taining less than 0.02% of higher hydro-Ba (OH)<sub>2</sub> solution and over ascarite to remove an impurity of CO<sub>2</sub> (0.7 vol %). NH<sub>3</sub> on methane reforming, ammonia decompowas used unpurified.

The methane reforming experiments were evaluated in a similar way as the ethane reforming experiments assuming first order kinetics with respect to methane as generally agreed in the literature (9, 10):

$$r = k \cdot p_{\text{CH},(1 - Q/K_p)},$$
 (7)

where Q and K, are reaction quotient and equilibrium constant for:

$$CH_4 + 2H_2O = CO_2 + 4H_2$$
. (8)

rates calculated at the same standard con-The activities were compared by the initial ditions as above with H<sub>2</sub>O/CH<sub>4</sub> = 4.

In the experiments on decomposition of ammonia:

$$2NH_1 = N_1 + 3H_2,$$
 (9)

the unconverted ammonia was absorbed in a solution of sulfuric acid and the conversion was calculated from the amount of washed exit gas and the ammonia feed.

The activity was expressed as the conversion at 500°C using a space velocity of 90 mol NH3/g cat/hr.

ethane on The experiments genolysis

$$C_2H_6+H_2=2CH_4$$

were performed with following feed:

The heated length of the reactor was 60 on, whereas the height of the catalyst bed was 10 cm. The catalyst zone consisted of 025-approx 10 g of catalyst as 1-2 mm particles diluted to constant volume with

heated furnace with 3 independent sections.

$$C_2H_6$$
 5 vol %  
 $H_2$  20 vol %  
 $N_3$  75 vol %.

whereas the order with respect to H2, man than 10%. Therefore, the activity was pressed as the first order rate constant vary. However, the partial pressure of can be considered as constant due to large excess and the conversions being Reaction (10) is generally reported first order with respect to C2Hs 300°C

by Schoubye (12). A mixture of H<sub>2</sub> (containing approx 1 vol % CO was used (8) teed (0.1 Nm³/hr). The reaction: The methanation experiments were in

initie control valve. The flows of water and lighted hydrocarbons were indicated on

burets, whereas the gas feeds were measured flowmeters. The axial temperature gradient in the catalyst bed was found to be approx 1°C/cm, whereas the temperature difference between the external tube wall

$$CO + 3H_2 = CH_4 + H_1O$$
,

and the axis of the bed varied between 2 and 10°C. On this basis the experiments

cannot be considered as strictly isothermal,

and the results may only be used for rela-Feedstock. The compositions of the feed-

ive comparisons of the catalysts.

% CO in H<sub>2</sub>; temperature = 250°C; pro ture in the range 200-300°C depending of 0.5 g. The initial rate was calculated the basis of the kinetics of Schoubye was performed isothermally at a temper the expected activity. The catalyst used as 0.4-0.5 mm particles in amo at the following standard conditions: 1 sure = 1.0 atm abs.

### Reforming Experiments at Elevated Pressure es;

gere desulfurized in a separate apparatus ver NiMo catalyst and ZnO. Analyses lowed contents less than 0.05 ppm S. The gaseous hydrocarbons, N2 and H2 were used

graphic analysis. The liquid hydrocarbons

štocks were determined by gas chromato-

through a vessel for mixing and a preheater fore the reactor. The reactor effluent was were pumped to an evaporator and mixed to a tubular reactor. H2 was added just be with gaseous hydrocarbons and N2 to be ditions. Water and liquid hydrocarbon fee used as feed. The feed mixture was pass depressurized and passed through a cool Apparatus. The apparatus was design to simulate operation under industrial 🥳 and a separator to a gas meter.

could not be maintained for more than a few hours. This could be ascribed to sulfur poisoning caused by the quick accumulation of sulfur at the extremely high throughputs. cluded only one measurement. As standard Therefore, most of the experiments inconditions were used: ameter of 10.0 and 17.5 mm, respectively. inm was placed in the axis of the reactor, and another at the external surface. The reactor was surrounded by an electrically The reactor was made of 18/8 stainless esteel and had an internal and external di-

carbon were taken and analyzed by means of a gas chromatograph and Orsat analysis sure was changed, this was done in an unsystematic way to minimize the influence of poisoning. Samples of dry exit gas and in some cases a sample of unconverted hydro-In a few experiments where the total pres-

> gradients were of the same order of size in all experiments. Above the catalyst zone

magnesium aluminum spinel. The temperafor was adjusted by thyristor controllers, and the pressure was maintained by a pneu-

The amount of catalyst was determined such that conversions and temperature

particles of magnesium aluminum spinel.

the analysis and flow measurement of the CH, formed, and from the measured consumption of liquid hydrocarbon giving the total g atoms of C in the feed. In the experiments with gaseous feed the conversion Assuming pseudo-first order kinetics with respect to the hydrocarbon an apparent higher hydrocarbons was calculated from exit gas giving the g atoms of CO, CO2 and Evaluation. The conversion, x, of the was calculated on basis of the exit gas only. constant ka was calculated from:

$$k_a = \frac{F}{W} \ln \left( \frac{1}{1 - x} \right). \tag{12}$$

situations where the kinetic expression is unknown, it may be useful for comparing space velocity is varied. However, it must be emphasized that the apparent rate conlyst performance at other conditions. It In an experiment with CH, as feed, x was replaced by  $x/x_{\rm eq}$  where  $x_{\rm eq}$  is the conversion to establish the equilibrium. As described by Hougen and Watson (13) for different catalysts to assume that even complex catalytic systems approximate to a pseudo-first order relationship if only stant obtained by this empirical method may hardly be used for prediction of catashould be used only for relative comparison

> apparatus was checked for leakages in H2 at 36 atm abs. The reactor was heated up ing H<sub>2</sub> to about 400°C. Steam and then

Procedure. Before each experiment the

\*unpurified.

experiments were performed with addition

the process conditions were established. All  ${}^{*}_{0}$  $H_{2}$ ,  $(H_{2}O/H_{2}=10)$ , to avoid oxidation. it soon turned out that the catalyst activity

gradually hydrocarbon feed were added and

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TABLE 2
PROPERTIES OF CATALYSTS: A-TYPES

							Partic	ele size	Activity reformi	for C <sub>2</sub> H <sub>6</sub> ng 500°C
Catalyst no.	Comments	Na (wt %)	Contents of red. Ni (wt %)	Ni area (m²/g)	Total area (BET) (m²/g)	n <sub>0</sub> N <sub>2</sub> -capacity (B <sub>b</sub> ) (N cm²/kg)	Calc.  r <sub>mean</sub> (Å)	Electron microscope particle φ (Å)	r <sub>i</sub> (mol/g/hr × 10)	r <sub>s</sub> (mol/ m² Ni/hr × 10³)
Group 1. Fi	ixed composition (25 wt % Ni)	. Na: 0.04-0	.16 wt %. Pre	eparation re	ute 1.					
A1	-	0.07	23.8	2.00	20.0	14	398	250-2500	2.4	120
Å15		0.04	22.0	3.01		34	245		4.40	145
A3		0.05	16.4	6.97	45.5	172	79	50-250	8.25	118
A4	•	0.05	23.1	3.78	33.3	5	205	500-1000	2.0	53
A16		0.14	22.9	2.99			257		1.5	52
	A16 sintered	0.16	~ 23.5	1.06			740	150-2000	0.51	48
A17		0.10	26.4	2.72		80	326		1.4	52
A18		0.08	25.7	3.46	27.4	50	249		4.2	122
Group 2. F	ixed composition (25 wt % Ni)	). Na < 0.01	wt %.~							
	Various preparation routes.									
. A19	Al washed in H <sub>2</sub> O					108			4.0	198
A20	Preparation route 2		18.1	3.71	15.7	112	163		8.2	222
A21	A20 sintered		10.1	1.09	2.34		311		2.0	183
A22 .	Preparation route 1		20.0	4.16	25.1	79	161		11.9	287
A23	A22 sintered		. 22.9	1.08	8.6	5	684		2.9	282
A24	Preparation route 1	Mr. Colomber 1 de	20.4	8.87	53.7	Contradence w	77	(a)	20.4	230
A25	Preparation route 3 2		E 20:3	6:34	39 8	79	107.	A PARTY	10.5	165
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Group 3. N	i content varied. Na < 0.01 w	t %. Prepara		4 17		96	112	50-125	10 6	254
Group 3. N A10	i content varied. Na < 0.01 w Ni/Mg = 1/1	t %. Prepare 0.04	13.9	4.17		96	112	50-125 75-200	10.6	254 90
Group 3. N A10 A26	i content varied. Na $< 0.01$ w Ni/Mg = $1/1$ 0.25/1	t %. Prepara	13.9 4.3	4.17 2.12	N. T.	96	68	75-200	1.9	90
Group 3. N A10 A26 A27	Ni/Mg = 1/1 0.25/1 0.5/1	t %. Prepara	13.9 4.3 8.2	4.17 2.12 4.35		96	68 63	75–200 75–200	$\begin{array}{c} 1.9 \\ 9.5 \end{array}$	90 212
Group 3. N A10 A26 A27 A28	Ni/Mg = 1/1 0.25/1 0.5/1 1/1	t %. Prepara	13.9 4.3 8.2 14.2	4.17 2.12 4.35 6.10		96	68 63 78	75–200 75–200 75–200	1.9 9.5 18.4	90 212 299
Group 3. N A10 A26 A27 A28 A29	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1	t %. Prepara	13.9 4.3 8.2 14.2 23.9	4.17 2.12 4.35 6.10 8.50		96	68 63 78 94	75–200 75–200 75–200 75–300	1.9 9.5 18.4 16.8	90 212 299 197
Group 3. N A10 A26 A27 A28 A29 A30	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1	t %. Prepara	13.9 4.3 8.2 14.2 23.9 35.1	4.17 2.12 4.35 6.10 8.50 8.45		96	68 63 78 94 139	75–200 75–200 75–200 75–300 200–600	1.9 9.5 18.4 16.8 18.6	90 212 299 197 221
Group 3. N A10 A26 A27 A28 A29	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1	t %. Prepara	13.9 4.3 8.2 14.2 23.9	4.17 2.12 4.35 6.10 8.50		96	68 63 78 94	75–200 75–200 75–200 75–300	1.9 9.5 18.4 16.8	90 212 299 197
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1	t %. Prepare	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35	eparation r		68 63 78 94 139 385	75-200 75-200 75-200 75-300 200-600 500-1000	1.9 9.5 18.4 16.8 18.6 8.0	90 212 299 197 221 219
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m	t %. Prepare	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35	eparation r		68 63 78 94 139 385	75-200 75-200 75-200 75-300 200-600 500-1000	1.9 9.5 18.4 16.8 18.6 8.0	90 212 299 197 221 219
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m	t %. Prepare	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23)	eparation r		68 63 78 94 139 385 485	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1	90 212 299 197 221 219 214
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m	t %. Prepare	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35	eparation r		68 63 78 94 139 385	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1	90 212 299 197 221 219 214
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48	eparation r		68 63 78 94 139 385 485	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1	90 212 299 197 221 219 214
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1 ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co Ni/Co = 3/1 romoted with alkali (25 wt %	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48	eparation r		68 63 78 94 139 385 485	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1	90 212 299 197 221 219 214
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33 Group 5. P	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co Ni/Co = 3/1  romoted with alkali (25 wt % A1 0.14 wt % K <sup>b</sup>	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48 6.34	eparation r		68 63 78 94 139 385 485	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1 0.020 0.016 2.9	90 212 299 197 221 219 214 1.6 0.29
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33 Group 5. P A34 A35	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co Ni/Co = 3/1  romoted with alkali (25 wt % A1 0.14 wt % K A1 0.53 wt % K	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48 6.34	eparation r		68 63 78 94 139 385 485	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1 0.020 0.016 2.9	90 212 299 197 221 219 214 1.6 0.29 46
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33 Group 5. P A34 A35 A36	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co Ni/Co = 3/1  romoted with alkali (25 wt % A1 0.14 wt % K <sup>6</sup> A1 0.53 wt % K <sup>6</sup> A25 2.1 wt % K <sup>6</sup>	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48 6.34		oute 1.	68 63 78 94 139 385 485 140 132	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1 0.020 0.016 2.9  0.79 0.097	90 212 299 197 221 219 214 1.6 0.29 46
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33 Group 5. P A34 A35 A36 A37	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co Ni/Co = 3/1  romoted with alkali (25 wt % A1 0.14 wt % K <sup>b</sup> A25 2.1 wt % K <sup>c</sup> A1 0.61 wt % Na <sup>b</sup>	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48 6.34		oute 1.	68 63 78 94 139 385 485 140 132	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1 0.020 0.016 2.9  0.79 0.097 0.075	90 212 299 197 221 219 214 1.6 0.29 46 27 3.7 2.1
Group 3. N A10 A26 A27 A28 A29 A30 A31 A32 Group 4. O A13 A14 A33 Group 5. P A34 A35 A36	Ni/Mg = 1/1 0.25/1 0.5/1 1/1 2/1 4/1 6/1 8/1  ther metals (total content of m Ni/Cu = 0.7/1.3 Ni replaced by Co Ni/Co = 3/1  romoted with alkali (25 wt % A1 0.14 wt % K <sup>6</sup> A1 0.53 wt % K <sup>6</sup> A25 2.1 wt % K <sup>6</sup>	t %. Prepars 0.04  netals 25 wt	13.9 4.3 8.2 14.2 23.9 35.1 41.9 48.3 %) Na < 0.00 7.6 22.9a	4.17 2.12 4.35 6.10 8.50 8.45 3.64 3.35 1 wt %. Pro (1.23) 5.48 6.34		oute 1.	68 63 78 94 139 385 485 140 132	75-200 75-200 75-200 75-300 200-600 500-1000 2000-5000	1.9 9.5 18.4 16.8 18.6 8.0 7.1 0.020 0.016 2.9  0.79 0.097 0.075 0.32	90 212 299 197 221 219 214 1.6 0.29 46 27 3.7 2.1

<sup>&</sup>lt;sup>a</sup> Red. Co or Co + Ni.
<sup>b</sup> Alkali added after reduction.
<sup>c</sup> Alkali added during preparation.

TABLE 3-PROPERTIES OF CATALYSTS: B-E TYPES

	·						Par	ticle size	•	for C <sub>2</sub> H <sub>6</sub> ng 500°C	
Catalyst type	Support material	Comments	Contents of red. Ni (wt %)	Ni area (m²/g)	Total area (BET) (m²/g)	n <sub>0</sub> N <sub>2</sub> capacity (B <sub>5</sub> ) (ml/kg)	Calc r <sub>mean</sub> (Å)	Electron microscope particle $\phi$ (Å)	r; (mol/g/hr × 10)	r <sub>s</sub> (mol/ m² Ni/hr × 10 <sup>3</sup> )	JENS
Туре В. В	sased on MgAl <sub>2</sub> O <sub>4</sub>									-	- ' <del>'</del>
B1	MgAl <sub>2</sub> O <sub>4</sub>		8.0	0.44	4.19	12	607	500-2500	1.6	366	жc
<b>B</b> 5	MgAl <sub>2</sub> O <sub>4</sub>	1.57 wt % Ka	11.1	1.35			272	000 2000	0.029	2.1	· ĭ
B6 ·	$MgAl_2O_4$	0.52 wt % Ka	17.0						0.54	61	RU
.B4	MgAl₂O₄	1.53 wt % Ka	17.5	0.88			664		0.074	8.3	P
Туре С. В	sased on Al <sub>2</sub> O <sub>3</sub>										ROSTRUP-NIELSEN
C1	η-Al <sub>2</sub> O <sub>3</sub>		16.5	7.01		65	79	100-400	5.5	80	53
C6	η-Al <sub>2</sub> O <sub>3</sub> Cl	1.7 wt % Kb		6.34	•	29	10	100-400	1.8	29	S
C7	η-Al <sub>2</sub> O <sub>3</sub> Cl	5.8 wt % Kb		3.89		0.5			0.45	11	
C8	η-Al <sub>2</sub> O <sub>3</sub> Cl	1.3 wt % Nab	•	6.78		13			3.2	47	
C9	η-Al <sub>2</sub> O <sub>3</sub> Cl	1.1 wt % Cab		7.24	•	115			2.2	30	
C2.	γ-Al <sub>2</sub> O <sub>3</sub>		17.9	5.66	106	238	106		9.2	163	
C3	γ-Al <sub>2</sub> O <sub>3</sub>		20	3.62	71.1	196	185		8.2	227	
C10	γ-Al <sub>2</sub> O <sub>3</sub> C3	2.1 wt % Ka		4.19	76.4	· 13	160		1.9	46	
C4	$\gamma$ -Al <sub>2</sub> O <sub>3</sub>	red. at 500°C	44.3	11.31		586	131		34.3	304	
C5	α-Al <sub>2</sub> O <sub>3</sub>		6.3	1.39			121		2.0	141	
Type D. B	ased on various mater	ials									
D2	MgO		5.3	1.61			146				
D3	MgO, Al <sub>2</sub> O <sub>2</sub> , (MgAl	$_{2}O_{4}) Mg/Al = 0.5$	19.1	1.01	44		146		4.3	266	
D4 6	ZrO, L	The officer with the second second	16 2	2 58		0.5	210	744	17.4 0.067	120	وازيدين
			C. 200 11 12 1		· ·	<u> </u>			مح و برسر صور		77.5-
D5	N. B. Waller	1437656						A COLUMN	Marie San		
	ZrO <sub>2</sub> /γ-Al <sub>2</sub> O <sub>3</sub>	Zr/Al = 0.44	25.0	5.70	A About the same of the	The sail Said Said Said	147	militar Salasian	11.5	201	1.540
D6	Cr <sub>2</sub> O <sub>3</sub>	red. at 500°C	60.0	7.92			254		19.9	252	
D8	SiO <sub>2</sub>	0' (41 0	21.2	12.67	145	000	53		16.8	133	
D9	SiO <sub>2</sub> /Al <sub>2</sub> O <sub>3</sub>	Si/Al = 3	21.2	4.43		382	76		1.1	25	
D10	SiO <sub>2</sub> /Al <sub>2</sub> O <sub>3</sub>	$Si/Al = 2.1$ $3.9\% Na^{\circ}$	25	2.45	44		336		0.44	18	
D11 ·	SiO <sub>2</sub> /MgO	Si/Mg = 1.67	25	2.42	33		346		0.58	24	
D12	D11 with 0.49 wt %		25	2.13	10		394		0.097	4.5	
D13	TiO <sub>2</sub>	, 11	37.0	1.00	10		125		0.36	36	
D14	Carbon	•	14.4	(0.8)			120	500-1700	0.040	5	
D7	Cr <sub>2</sub> O <sub>3</sub> -	red. at 500°C	12.3	5.43			76		14.1.	260	
							•				
	ntaining precious met		0.5.	0.04-							Z
E1	Al <sub>2</sub> O <sub>3</sub>	Pt	0.5	0.24					0.53	221	Ni catalysts
E2	Al <sub>2</sub> O <sub>3</sub>	Pt	5.0	1.54					1.2	77	C <sub>A</sub>
E3 ' .	. carbon <sup>d</sup>	Pt	0.5	0.55					0.05	040	TΑ
E4 E5	Al <sub>2</sub> O <sub>3</sub>	Pd	0.5	0.41					0.87	212	ξ
E6	Al <sub>2</sub> O <sub>3</sub>	Pd Pd	5.0	3.46				•	6.7	180	ST
	γ-Al <sub>2</sub> O <sub>3</sub>		0.5	0.10 0.20					0.19	194	
E7 E8	carbon Al <sub>2</sub> O <sub>3</sub>	Pd Ru	0:5 0.5	0.26					5.4	2059	Õ
E9	Al <sub>2</sub> O <sub>2</sub>	Ru	5.0	1.81					5.4		æ
E10	$\gamma$ -Al <sub>2</sub> O <sub>3</sub>	Ru	5.0	0.55					30.3 0.20	1673	R.E.
E10	carbon <sup>d</sup>	Ru	0.5	0.48					0.20	37	Ođ
E12	γ-Al <sub>2</sub> O <sub>3</sub>	Rh	0.0	0.48					4.1	2502	RM
E13	carbon <sup>d</sup>	Rh	0.5	0.37					4.1	2002	FOR REFORMING
E14	γ-Al <sub>2</sub> O <sub>3</sub>	Re		0.21	•				0.10	50	Ф
		• • • • • • • • • • • • • • • • • • • •								<del></del>	

<sup>Added during preparation.
Added before reduction.
Ni area estimated from particle diameter.
Diluted with γ-Al<sub>2</sub>O<sub>2</sub>, 1/10.
Following figures based on metals indicated.</sup> 

of catalyst activities or reactivities of various hydrocarbons.

### 4. Catalysts

The catalysts which are listed in Tables 2 and 3 have been divided into various types determined by their chemical compositions. The A-type is based on magnesia containing approx 6 wt % Al. Group 1 has whereas groups 2-4 are free of alkali metals (less than 0.01 wt %). Group 2 includes catalysts with fixed composition but with content is varied in group 3. The B-type is whereas the C-type is supported by various types of alumina. Catalysts based on other no alkali metals (less than 0.01 wt %). The different preparation routes whereas the Ni based on magnesium aluminum spinel finally, type E covers catalysts containing precious metals. If not indicated in the tables the catalysts of groups B-E contain basis for comparisons. In addition to what is indicated in Tables 2 and 3 such groups supports are represented by Type D and numbering of the catalysts is made in accordance with the numbering used previously (14). Some of the catalysts were prepared in parallel and may present a better are (A20-A23), (A26-A32), (B1 and B4-B6), and (A25, A36, C7, C8, D8, D10 and a minor content of Na (0.02-0.16 wt %)

The catalysts were reduced in a separate reactor at 850°C in H<sub>2</sub> for a period of 2 hr as previously (14). The nickel surface area contain 44.2 × 10-9 g S/cm2. The sulfur unless otherwise indicated. The content of nickel in the reduced state was determined was calculated on basis of the sulfur capacity assuming the saturated surface to capacity was determined by chemisorption of H2S at 500°C as described previously (15). A mean particle radius was calculated from the nickel surface area and the concation. For some catalysts the range of nickel particle size was estimated by means of an electron microscope. A N2 capacity of This figure should be used only as an indithe Ni surface was determined by a procetent of reduced nickel as shown earlier (14) dure described below.

# - 5. Adsorption Studies

The presence of special sites on the nickel surface was investigated on basis of the work by Van Hardeveld et al. (16-19) who found that N<sub>2</sub> was physisorbed on the so-called B<sub>3</sub>-sites of nickel crystallites. In the range 15-70 Å in a polarized state which was infrared active. The polarization explains the relatively high initial heat of adsorption (approx 13 kcal/mol). Other studies (20-22) have considered the IR active is a weakly chemisorbed molecules whereas the results of Bradshaw and Pritchard (28, 24) appear consistent with those of Van Hardeveld et al.

As demonstrated by Van Hardeveld and sorbed at room temperature and relatively The results make it likely that all N<sub>2</sub>, ad adsorbed N2 and the number of B5-sit of the adsorbed volume at room tempë ture and a  $m N_2$  pressure of 200 mm Hg  $m a_2^2$ Van Montfort (19) the determination intensity of the IR-absorption band of Hardeveld (25) has proposed measurem the number of B<sub>s</sub>-sites either by IR or  $v_{
m c}$ metric methods is not unambiguous, there is no simple correlation between which can be adsorbed at higher pressure low pressure is adsorbed in the IR-act form, but this N<sub>2</sub> is only a fraction of t standard method for determination of lower temperature. On this basis number of B<sub>5</sub>-sites.

for approx 3 hr at 800°C to a pressure was heated in a flow of H2 and evacuated below 10-4 mm Hg before the measure was necessary to remove all H<sub>2</sub> from the surface. It was evident that H2 could still be removed when heating the evacuated sample from 600 to 800°C. Evacuation were performed in a conventional BET ap This method has been adopted here, an sorbed N2, the measurement was repeate umes at 200 mm Hg. The measurement The high evacuation temperation after poisoning the nickel surface with N sample of 15–20 g of prereduced cataly as a correction for ordinary physically The  $N_2$  capacity,  $n_3$ , was determined as difference between the two adsorbed v paratus as described previously (15) ments.

on the line N<sub>2</sub> capacity. Catalyst C4 was reduced basis of at 800°C but evacuated the second catalyst C6.

Some isotherms are shown in Fig. 2. For catalyst C4, which was the catalyst with the highest N2 capacity, the contribution incertain. Even catalyst C4 shows a value finde less than the typical values of the of the small adsorption volumes makes an from the physically adsorbed N<sub>2</sub> is negligible, whereas the N2 capacity of a typical catalyst like A1 is determined as the difference between two figures of same order. his makes the determination of  $n_0$  very of no of approximately one order of magnisplained by the larger nickel crystallites of the present catalysts. The high uncertainty estimation of the initial heat of adsorption very unsafe. On basis of the isotherms on 34 at 25 and 65°C an initial isosteric heat of adsorption was estimated to about 11 catalyst of Van Hardeveld. This kcal/mol N2.

### RESULTS

# Ethane Reforming

A series of experiments was performed to determine a kinetic expression for ethane reforming at 500°C over catalyst A1. Mini-

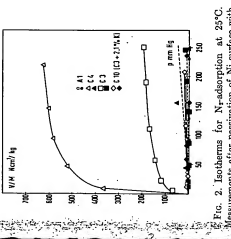


Fig. 2. Isotherms for N<sub>T</sub>-adsorption at 25°C. Measurements after passivation of Ni surface with No indicated by filled points and broken lines.

mum variance for an expression of the form (5) was obtained by:

 $r = 1.67 \times$ 

105 e-18100/RTpczH6.54pH10-0.33pH10.2 mol/g·hr.

The effect of the partial pressure of carbon dioxide appeared insignificant. Experiments at 450 and 550°C including variations of the partial pressure of steam showed  $\alpha_{\rm H_20} = -0.60$  and -0.23, indicating that a more complex expression is required to describe the kinetics of the reaction in a broad temperature range. In this connection it should be noticed that Bodrov, Apel baum and Temkin (9) found the kinetic coefficient of hydrogen to vary with temperature in their studies of the methane reforming reaction.

respect to steam varied significantly from showed negative values of  $\alpha_{\rm H_20}$  down to -0.5, whereas catalysts based on magnesium aluminum spinel (B1) or alumina tive, respectively. The addition of potas-The kinetic coefficients, mainly for steam, were determined for some other catalysts. As shown in Table 4 the reaction order with catalyst to catalyst. Thus, catalysts containing free magnesia (A-types and D3) (C1) showed values of zero or slightly posisium caused a significant decrease of an,o. The addition of sodium had a less pronounced effect. The addition of potassium implied no change of the apparent activation energy whereas replacement of nickel with a nickel-copper alloy caused a substantial increase of the activation energy. Phis is demonstrated in Fig. 3.

The influence on the experiments of mass and heat transfer was analyzed by means of the Topsøe REACTOR program (26) designed for computer calculation of the temperature and conversion profiles of a fixed bed converter. The calculation is performed on the basis of the intrinsic kinetics and the pore volume distribution of the catalyst. For catalyst A18, a catalyst at a typical activity level, the calculation showed an effectiveness factor of more than 0.95 and a temperature drop over the gas film surrounding the particles of 1.3°C, which reflects only negligible restrictions. For a very active catalyst such as A22,

ETHANE REFORMING AT ATMOSPHERIC PRESSURE, 500°C: SUMMARY OF KINETIC PARAMETERS TABLE 4

è go

Catalyet		. Ki	Kinetic coefficients	g		
type Comments		C2H,	Н,0	H <sub>2</sub>	- Apparent activation energy $E_a$ , (kcal/mol	Apparent activation mergy $E_a$ , (kcal/mol
Type A, Group 1 (MgO) A1 A18		0.54	-0.334 -0.52b	0.2	18.1	(0.15)
. Group 2						
		-	0.0			
. A21 A22			- 0 .26 - 0 .48			
A23	,		-0.17			genden ve digi
A13					24.4	(0.7)
Group 5 (Promoted) A34 A1, 0.14 wt % K	oted)		-0 4%		18 0	6 0
A35 A1, 0.53 wt % K A37 A1, 0.61 wt % Na			-1.81• -0.7	•	19.6	(0.5)
Type B, (MgAl <sub>2</sub> O <sub>4</sub> )						100
B1 B4 1.53 wt % K	14	(0.4)	0.06		18.3	(2.5)
Type C, (Al <sub>2</sub> O <sub>2</sub> ) C1		(0.6)°	0.13			
Type D			6			TO THE
$D_4 \qquad ZrO_1$	. 1/1		-0.26		19.2	(0.9)

5.ريب **.**.. I Figures in brackets indicate the accuracies of the activation energies. The accuracies of the coefficie are estimated to:  $^{a} \le 0.05$ ;  $^{b} 0.05-0.1$ ;  $^{c} 0.1-0.2$ ;  $^{d} 0.2-0.3$ ;  $^{o} 0.3-0.5$ .

some transport restrictions were indicated by an effectiveness factor of approx 0.9 and a temperature drop over the gas film of

Due to the variations in kinetic coeffirates ri as described previously. For cataysts for which no kinetic coefficients were determined, r, was calculated on the basis of the kinetics of a similar catalyst. The 10%, respectively, which yield a relative cients from catalyst to catalyst the activities were compared on the basis of initial relative standard deviations of  $r_i$  calculated from experiments based on samples of the same origin and of the corresponding nickel areas were estimated to be approx 17 and standard deviation of the specific rate r, of no better than 20%. Therefore, the experiments only allow identification of substantial differences of specific activity.

From the results shown in Tables 2 and 3

0.1-0.35 mol/m<sup>2</sup> Ni/hr. This is observed on a broad range of support materials such as magnesia in various sintered forms, and with AI/Mg ratios varying from 0 to 2 magnesium aluminum spinel, alumina in different modifications, alumina stabilized by zirconia, chromia and silica. Moreover, results from a systematic series of catalysis (A22-A32) shown in Fig. 4 indicate only insignificant influence on the specific achave a specific activity in the range re tivity of the nickel content and of the nickel crystallite size varying with the ments with catalysts with nickel crystallig the alkali-free nickel catalysts appear the smallest crystallites but more exper below 70 Å are required to elucidate 🚓 results from a systematic series of cataly nickel content from 50 to 5000 A. The sults might reflect less specific activity,

Some alkali-free catalysts show a small

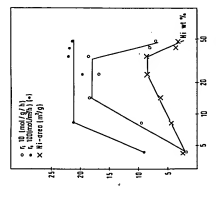


Fig. 4. Influence of Ni content on Ni surface area and activity for ethane reforming at 500°C. Catalysts A26-A32.

the specific activity by only approx 50%. chlorine and arsenic being less than that of This appears from Table 5 showing in addition an influence of other poisons such as tassium is remarkable as sulfur poisoning of catalyst A1 involving a gradual blocking of the nickel surface is causing a decline of potassium.

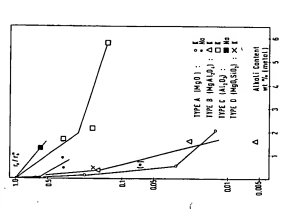
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Fig. 3. Temperature dependence of rate constant

Of ethane reforming at 500°C.



· Fig. 5. Influence of alkali on the specific activity for ethane reforming at 500°C.

on specific activity. The catalysts based on itania had  $r_s = 0.02-0.04 \text{ mol/m}^2 \text{ Ni/g}$ , whereas the activity of catalysts based on with  $r_s = 2.6 \times 10^{-3}$  and  $3.8 \times 10^{-4}$ , respecjeing supported mainly by alumina as it jure zirconia and carbon was very poor lively. The low activity of the zirconia based catalyst (D4) is not accompanied by Dased causayou (127) and activation energy differing from that of alumina based catalyst (D5) appears norcatalyst A1. The activity of the zirconiamal, which may be explained by nickel The addition of potassium to the cata-Fas present as nickel aluminum spinel silica-alumina, silica-magnesia and before the reduction.

potassium. The significant influence of poysts can result in a decrease of the specific fect of sodium addition is less than that of activity of more than one order of magniude. This is demonstrated in Fig. 5 which indicated the influence to be stronger on 4- and B-types than on C-types. The ef-

THE INFLUENCE OF POISONS ON THE ACTIVITY OF CATALYST AI TABLE 5

Expt no.	Poison	Content (wt ppm)	Sulfur capacity (wt ppm).	Coverage	$r_i \times 10$ (g mol/g/hr)	r. × 10 (g mol/ m² Ni/hr
Av of unpoisoned A1	1	80	883	<0.1	2.41	120
4201	Ω	239	805	0.30	99.0	62
4202	Ø	360	805	0.45	0.53	69
4203	Ø	398	805	0.49	0.59	. 79
329	Ø	615	805	0.76	0.38	56
55	ß	802	805	1.00	<0.01	Î
133	ប	1350	882	۵.	2.43	7
141	As	4200	885	٥.	0.55	all a series

mined by chemisorption of hydrogen sulfide The presence of potassium implied no shown in Tables 2 and 3 were significantly the nitrogen capacity of the zirconia-based catalyst D4 was negligible, and nitrogen capacity. This has been done decrease of the nickel surface area deterat 550°C or as demonstrated previously -72°C. However, the measurements of nitrogen adsorption on the nickel surface it is natural to correlate catalyst activity in Fig. 6 and a correlation coefficient of 0.94 indicating great significance was calchemisorption of hydrogen at influenced by the addition of potassium. As, moreover,

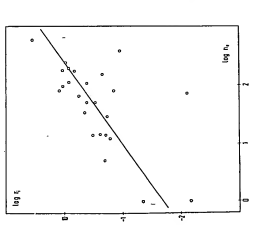


Fig. 6. N<sub>2</sub> capacity and activity for ethane reforming at 500°C.

measurements and does not account for the relatively low activity of the silica-alumi supported catalyst D9, it seems likely th the great uncertainty of the adsorpt face. It does not necessarily ascribe a st cial high activity to B<sub>5</sub>-sites, as the nuṃ culated. Although the correlation refl the differences in specific activities are such as corner atoms with low coordinat of these sites in some way is correlated w the number of other surface configurat lated to inhomogeneities of the nickel numbers.

specific activities of metals based on alum Measurements on catalysts contain ina or magnesia may be listed as follows different metals showed large variations specific activities. From Tables 2 and 3

The low activity of cobalt may be related by some irregularities in the preparati The low activity of E10 could be explain oxidation of cobalt by steam. The preci ratio close to the equilibrium constant. metals supported by carbon showed Rh, Ru > Ni, Pd, Pt, > Re > (Nio. Cu1. 1) > to the process conditions with a  ${
m H_2O}$ poor activities similar to the results carbon supported nickel catalysts.

# 2. Other Reactions at Atmospheric Pressure

applied for methane reforming experiments This was justified by an experiment show catalyst A1, ethane was replaced by meth ane or n-butane. First order kinetics we some reforming experiments

REFORMING EXPERIMENTS AT ATMOSPHERIC PRESSURE VARIOUS HYDROCARBONS, CATALYST A1 TABLE 6

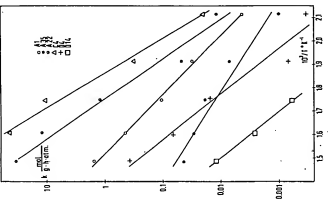
A on a optive tion	$\begin{array}{ccc} \operatorname{ch} & \operatorname{ch}$	(kcal/mol)	26.2 (0.4)	18.1 (0.15)	18.6 (0.6)
Sp act (500°C)• × 10°	rec (g atom/	m² hr)	. 61	240	552
Sp act (	r, (mol/	m² hr)	19	120	138
	runde für	Feed	CH,	C,H,	" "C,H10

 Rates calculated for same partial pressure of  $H_2O/H_2 = 10;$  $(H_2O/C_nH_m = 8;$ ndrocarbon 50°C)

A Figures in parentheses indicate the accuracy of tion of butane reforming experiments. The results are summarized in Table 6. The ing the kinetic coefficient to be close to carbon. The apparent activation energy is higher for methane than for ethane reformon atom is higher for n-butane than for thane. Moreover, it was observed that in tion with n-butane resulted in no higher initial specific rates have been calculated at the same partial pressure of the hydroing, and a smaller reactivity is reflected by lower initial specific rate. Contrary to the temperature range 400-525°C the reacunity. Equation (13) was used for evaluahis n-butane shows an activation energy and molar specific rate very close to that of othane. This implies that the rate per car-

by calculating activities relative to catalyst Results from experiments on hydrogenolmonia are shown in Figs. 7 and 8, respeclively. The trends of activities for these is of ethane and decomposition of amreactions, methane reforming and methanation of carbon monoxide were compared with the activity trend for ethane reforming A1. The results are shown in Table 7. ydrocarbons among the products.

The data for methanation show some deviane reforming, ethane hydrogenolysis and methanation are broadly in line. Thus, the The activity trends for ethane and methpresence of potassium affects all these reactions by decreasing the specific activity.



Temperature dependence of rate constants for various catalysts. Fig. 7. Ethane hydrogenolysis.

Firstly, the catalysts based on zirconia (D4) and carbon (D14) show relatively high activities. Secondly, it appears that the ethane reforming. On the other hand, the influence of alkali appears to be more proparticularly those based on alumina, is higher than the corresponding values for ations from those for ethane reforming. specific activity of highly active catalysts, nounced for methanation.

catalysts and D4. No large changes of specific activities are observed, and as shown position correlate simply with the nickel The results from decomposition of ammonia differ significantly from the general as no decrease of the specific activity is observed for potassium-promoted in Fig. 9 the activities for ammonia decomarea. The cobalt-containing catalyst (A33) fits into this correlation. trend.

# 3. Reforming Experiments at Pressure

The reforming experiments at pressure with 1-2 mm particles were subject to great

Decomp. of NH<sub>1</sub> relative conversions x<sub>e</sub>/ x<sub>e</sub>[A1]

ä

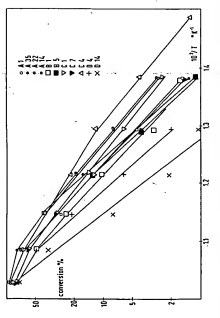


Fig. 8. Ammonia decomposition. Temperature dependence of conversion at standard conditions.

creases with pressure for catalysts Al and for the strongly alkalized B4. These results

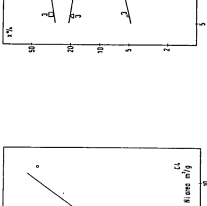
B6, whereas the opposite effect is observed

uncertainty as explained previously. This makes a more detailed evaluation doubtful. However, the results plotted in Fig. 10 show the conversion of ethane to increase with results from the experiment, 1231, perserved depending on the type of catalyst. pressure. In principle, this could be explained by Knudsen diffusion causing severe restrictions at low pressure but the formed at constant pressure and varying nitrogen flow indicates the pressure effect on the conversion to be related to the kinetics. When using naphtha as feed, posi-As shown in Fig. 11 the conversion intive and negative pressure effects are ob-

ethane reforming at atmospheric pressure The activities of various catalysts obatmospheric pressure. This is demonstrated in Fig. 12, where the straight line reflects listed in Table 4 which indicates the overall pressure coefficient to be positive for A1 and abs correlate with the corresponding activit ties determined by ethane reforming at tained by reforming of naphtha at 31 atm the correlation for equal specific activities correlate with the kinetic coefficients for B6 and negative for B4.

RELATIVE SPECIFIC ACTIVITIES AT ATMOSPHERIC PRESSURE TABLE 7

		Reforming	ming	Hydrogen-	Methana-	
	Catalyst	1 7	120	- olysis of	tion of	Decomp
		ا چ آ	Ţ C	C,H	3	NHS
No.	Remarks	200°C	200°C	300°C	250°C	200°C
A1	(0.07 wt % Na)	1.0	1.0	1.0	1.0	1.0
A22		2.4	2.0	4.2	9.1	6.0
A35	0.53% K	0.03	0.09	0.08	0.002	6.0
Bl		3.0	1.4		7	3.2
B5	1.57% K	0.02	0.02			1.5
<u>ವ</u>		0.7			2.7	0.4
2		2.5		8.7	20.6	0.5
3	5.8 wt % K	0.09			0.001	0.5
D4	ZrO,	0.03	0.01	0.08	8.0	0.4
D8	SiO <sub>2</sub>	1.1			4.9	in full
D3	SiO,/Al,O,	0.2			0.04	Ť
D14	Carbon	0.04	0.04	0.01	0.4	0.5



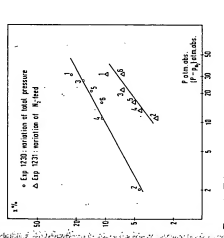
Fra. 9. Ammonia decomposition at 500°C. Acdivity as function of nickel surface area. Activity expressed as conversion at standard conditions relative to catalyst A1.

5

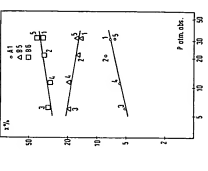
A 14

A22

Experiments with various hydrocarbons performed at 31 atm abs and 500°C using a fixed steam to carbon ratio revealed great differences in conversions. As shown in Table 8 most hydrocarbons are more reacanaphthenes whereas the reactivity of benzene is very close to that of methane. Howpear less reactive than isoparaffins and tive than is methane. Normal paraffins apever, it should be noticed that the experi-



Rependence of conversion at standard conditions at \$500°C. 0.5 g catalyst A18. The numbers indicate Fig. 10. Ethane reforming at pressure. Pressure the sequence of measurements.



conditions at 500°C. Expt 1206: 0.5 g Al; Expt 1195: 10.7 g B5 (1.5 wt % K); Expt 1196: 11.7 g Pressure dependence of conversion at standard B6 (0.3 wt % K). Numbers indicate sequence of Fig. 11. Reforming of naphtha 36 at pressure. measurements.

pressure of the hydrocarbons varies with ments are performed at a fixed steam to the number of carbon atoms of the hydrocarbon. As mentioned previously, extrapthe apparent rate carbon ratio which implies that the partial constant  $k_a$  is very doubtful. When the conby means of olation

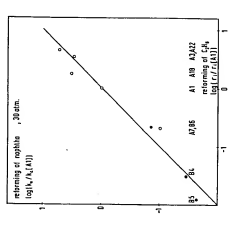


Fig. 12. Correlation between relative activities at 500°C for reforming of naphtha 36 at 30 atm and reforming of ethane at atmospheric pressure. Activities expressed relative to catalyst A1.

TABLE 8 REFORMING AT PRESSURE

Exp.	Feed	H <sub>2</sub> O/C (mol/atom)	p <sub>саны</sub> (atm)	Space velocity g atoms $\cdot$ 22.4 kg cat. $\times$ 10 <sup>3</sup>	Conversion to CO, CO <sub>2</sub> , CH <sub>2</sub>	k <sub>a</sub> calc. from (13) (g atom/g/hr)	$k_a \times$ (no. C-atoms/ no. C-C bonds)	$k_a  imes rac{p_{ ext{CH}_i}}{p_{ ext{C}_n  ext{H}_m}}$
1211	Methane	3.71	6.0	194	4.65b	0.43		0.43
1212	Ethane	3.79	3,3	207	31.5	3.5	1.8	6.4
1230		3.85	3.2	204	22.5	2.3	1.2	4.3
1216	(n,iso)Butane)	3.85	1.7	204	39.9	4.6	3.5	16
1210	Cyclohexane	3.73	1.2	209	53.9	7.2	7.2	36
1208	Benzene	3.96	1.1	200	5.7	0.53	0.53	2.9
1209	<i>n</i> -Heptane	3.83	1.0	206	18.7	1.9	1.6	11
1214	Trimethyl-butane	3.88	1.0	201	33.2	3.6	3.1	22
1213	n-Decane	3.77	0.7	104	32.8	1.9	1.7	16
1203	Naphtha 36 (FBP 120°C)	3.74		209	40.0	4.8	_	_
1215		3.76	_	209	35.8	4.1		
1207	Naphtha 49 (FBP 169°C)	3.69		212	15.5	1.6		

Experiments with various hydrocarbons; Catalyst A18 (0.5 g as 1-2 mm particles); temp. 500°C; pressure 31 atm abs; H<sub>2</sub>O/H<sub>2</sub> = 10.

<sup>b</sup> Conversion to CO and CO<sub>2</sub>.

Wing appear complex it may be helpful to giquence. Ethane is assumed to be chemither respectively, the following sequence can sorption has also been suggested in the litalkali. A rôle of the carrier for steam adgorbed on nickel following the pattern generally proposed in studies of ethane discuss qualitatively the parameters inerature (4, 6, 8). When  $S_1$  and  $S_2$  are empty yolved in terms of a simple speculative sesupport material and of the presence of kinetic coefficient for water of the type of carbon-carbon bond and formation of a hydrogenolysis (11). An initial chemisorpsites on the surface of nickel and the carthe basis of the observed influence on the on the carrier is assumed to be involved on surface radical CH<sub>2</sub>. Adsorption of steam genation is followed by a rupture of the tion step on a dual site involving dehydro-Although the kinetics of ethane reformand the chemisorption of methane to reaction with steam being the slow step. limiting at high temperature. to be rate determining at low temperature reforming assumed the reaction with steam the type of hydrocarbon, whereas Bodrov, Apel'baum and Temkin (9) for methane that the rate determining step varies with Phillips, Mulhall and Turner (7) suggested favor of the desorption of products or the and Dixon (5) and Saito et al. (8) were in the rate determining step, whereas Bhatta the rupture of the carbon-carbon bond as Slovokhotova and Balandin (4) argued for kinetics for reforming reactions. Balashova, assumptions made in the formulation of

ered as irreversible since no influence on the surface reaction (18) might also be considstudy. Secondly, for practical purposes, the in the temperature range of tion of ethane is most probably irreversible Boudart (34), the dissociative chemisorptermining step. Firstly, as proposed by the kinetics by means of a single rate de-There are some objections to discussing

 $C_2H_6 + 2S_1 = (S_1)_2 - C_2H_z + \frac{6-z}{2}H_2$ , (14)

cordance with the results obtained at low spect to the hydrocarbon, and it is in acatoms. This trend is still obtained when assuming a reaction order of 0.5 with reresults indicate normal paraffins to be more stants are corrected for the actual partial pressure shown in Table 5. reactive with increasing number of carbon first order kinetics as shown in Table 8, the pressures of the hydrocarbons and assuming

a detailed evaluation of the analysis of the appears very complicated. unconverted naphtha from experiment 1207 yery unlikely. The experiment with benzene ence of intermediates in the products is in accordance with the result obtained in represents an exception to this picture, and atmospheric pressure. Therefore, the presthe reforming experiment with n-butane at cate the composition to be very close to that of the feedstock. This observation is the reactor effluent shown in Table 9 indi-The analyses of higher hydrocarbons in

 $(S_1)_2-C_2H_z + H_2 = 2S_1-CH_z,$ 

 $S_2-H_2O + S_1 = S_1-O + H_2S_2$  $H_2O + S_2 = S_2 - H_2O$ ,

 $S_1-CH_x + S_1-O = 2S_1 + CO + \frac{x}{2}H_2$ 

(18)

(17)(16)(15)

 $CO + S_1 - O = S_1 + CO_2$ (19)

may be converted to carbon. available for the reaction. Eventually they simplification because there may be a gradand may therefore diminish the surface Certainly, steps (14) and (15) represent a by Kemball (27) and Frennet and Lienard by further dehydrogenation. As discussed ual formation of strongly adsorbed species (28) these species may become less reactive

reaction as two irreversible steps. studies (31-33) it is believed that the dein quasi-equilibrium (11, 29, 30). In other sorption of products is rate determining. Recently Boudart (34) has considered the the overall reaction, all other steps being sumed to be the rate determining step of the carbon-carbon bond is generally as-For ethane hydrogenolysis the rupture of These opposite views are reflected by the

Discussion

JENS R. ROSTRUP-NIELSEN

TABLE 9
REFORMING EXPERIMENTS WITH VARIOUS HYDROCARBONS<sup>a</sup>

<b>7</b>		16 ane		10 nexane		08 zene	12 n-Hej		1214		1215 1203	12 Napht	
Expt no.: Feed: Analyses (wt %)	Feed	Liq. prod.	Feed	Liq. prod.	Feed	Liq. prod.	Feed	Liq. prod.	- Tri meth. buta. Feed	1213 n-Decane Feed	Naphtha 36 <sup>d</sup> Feed	Feed	Liq. prod.
C <sub>2</sub> H <sub>4</sub> C <sub>2</sub> H <sub>6</sub> C <sub>3</sub> H <sub>8</sub> + (C <sub>2</sub> H <sub>6</sub> )	0.01 0.05 1.20	0.04 0.47 1.57		,								0.10 1.13	
n-C <sub>4</sub> i-C <sub>4</sub> C <sub>4</sub> H <sub>8</sub>	80.30 17.92 0.27	79.00 18.61 0.00							0.05		0.02	3.70 1.08	
n-С <sub>5</sub> i-С <sub>6</sub> СР	0.25	0.31			0.01	0.01	0.06 0.01	0.02	$\begin{array}{c} 0.32 \\ 0.11 \end{array}$		29.16 8.55 1.74	3.92 4.45 0.51	2.05 0.22 0.22
n-C <sub>6</sub> i-C <sub>6</sub> MCP(+22DMP) CH(+33DMP) Benzene			99.98	1.65 0.02 0.92 96.34	0.01 0.04 0.07 99.71	0.08 0.04 0.03 0.06 62.23	0.82 0.06 0.17 0.47 0.05	0.57 0.04 0.10 0.36 0.24			16.51 16.84 3.40 1.86 1.24	4.39 4.70 2.72 1.80 0.23	4.51 3.58 1.20 1.19 0.39

n-C <sub>7</sub> 180-C <sub>7</sub>	0.01	0.72	0.02	0.38		87.71	0.01		5.14	5.21	7.02	
DMCP		0.18	0.07	0.17 0.11	$8.45 \\ 1.72$	7.67	98.78		5.45	4.78	5.50	
MCH(+22DMH + 113TMCP)	0.01	0.01		0.35	0.72	$\frac{1.52}{0.69}$			1.53	4.31	3.06	
ECP(+25DMH)	0.01	0.00		0.33	0.72	0.09			1.72	6.66	5.49	
Toluene									0.52	1.04	1.15	
Totuene		0.03		0.38	0.37	0.36			1.76	1.82	2.70	
n-C <sub>8</sub>				3.26	0.31	0.35		•	1.02	6.40	7.16	
$iso-C_8+C_{8+}$		0.06	0.07	3.30	0.20	0.19			2.33	11.87	14.60	
ECH	-			0.82					0.09	2.53	1.74	
Ethylbenz				0.54					0.12	0.30	1.10	
Xylenes				3.00					0.33	4.36	4.48	
				0.00					0.55	4.00	4.40	
n-C <sub>9</sub>				5.06					0.37	4.60	5.02	
C <sub>9+</sub>				10.26					0.17	10.43	15.41	
							•					
n-C <sub>10</sub>				1.61				>95	0.06	0.18	1.43	
C <sub>10+</sub>				7.02					0.07	2.07	9.62	
$n$ - $C_{11}$				0.09							0.00	
C <sub>n+</sub>				1.09							0.06	
~n+				1.05							1.10	

<sup>&</sup>lt;sup>a</sup> Temp:  $500^{\circ}$ C; pressure: 30 atm abs; catalyst: A18; gas chromatographical analyses of feed and liquid products. <sup>b</sup> CH<sub>4</sub>: 0.01 vol %.

<sup>c</sup> Calculated from gas analysis.

d Naphtha no.:	36	49
ASTM: IBP/MBP/FBP (°C)	40 65 120	40 89 169
Spec gr (g/ml)	0.674	0.705

ably also S<sub>1</sub>-O may be assumed to be the most abundant surface intermediates, the concentrations of other intermediates being negligible. There is some evidence for this assumption. As no intermediates are found in the products when using n-butane or higher hydrocarbons (except benzene) as this may indicate the same active species, containing one carbon atom, to be involved in the reaction. In addition, some LEED studies by Maire et al. (35, 36) have rate of carbon dioxide was indicated. Moreto be formed by adsorption on nickel of over, the surface species  $S_1$ -CH<sub>x</sub> and probshown similar structures of the type CH<sub>2</sub> methane, ethane, propane and neopentane. feed,

These assumptions lead to the following the total number of sites, L and M, on the equations for the rates of the steps and for surface of nickel and carrier, respectively:

$$r_{14} = r_{15} = \frac{1}{2}r_{18} \quad r_{+16} = r_{-16} \quad r_{+17} = r_{-17},$$
  
 $(S_1) + (S_1 - CH_z) + (S_1 - O) = L,$   
 $(S_2) + (S_2 - H_2 O) = M,$  (2

ment of the equilibria (2) and (3) being kinetically insignificant in accordance with ing a homogeneous surface and no changes heats of adsorption with coverage the the remaining steps involved in establishthe principle formulated by Boudart (34). Using Langmuir equations and thus assumfollowing rate equation is obtained: oţ

$$r_{+17} = r_{-17}$$
Equation (21) may the power-rate law using the proximation of the term  $r_{+17} = r_{-17}$ 

$$r = k_n p_{\text{Ch1}^{1-2m}} \cdot p_{\text{H}_1\text{O}^{2(n-m)}} p_{\text{H}_2^{2(m-n)}},$$
 (3)  
where  $0 < m < 1$ , and  $0 < m < 1$ .  
This expression may explain the kine coefficients obtained for catalyst A1 shown in (13), and that the kinetic coefining for steam may become less than a sheeved for catalysts A35 and B4 has a cheeved for catalysts A35 and B4

$$= \frac{k_A L^2 \cdot p_{\text{CiH4}}}{[(1 + (2k_A/k_7) (1/K_w)(p_{\text{CiH4}}/p_{\text{Hi}})p_{\text{Hi}} + K_w(p_{\text{Hi}}o/p_{\text{Hi}})]^2},$$
(21)

where  $k_A = k_{14}$ ,  $k_r = k_{18}$  and  $k_w = k_{+18}$ . k+17/(k-18·k-17).

tively to comprehend the varying kinetic coefficients reported in the literature (Table steam adsorption,  $K_{\omega}$ , and the relative The kinetic order with respect to steam may become positive or negative depending carbon adsorption,  $k_A$ , and the surface reaction,  $k_r$ . Since  $K_w$  is the product of the sizes of the rate constants for the hydro-From this expression it is possible qualita-1) and observed in this study (Table 4) on the size of the equilibrium constant for equilibrium constants of steps (16) and (17) it is strongly influenced by the

action orders is clear since the relative size ferent reactivities of various hydrocarbons sorption properties of the carrier material of the terms of denominator may change the activation energies for the hydrocarbon adsorption and the surface reaction, and the heat of adsorption for steam. The dif A possible temperature dependence of resecond term of the denominator in (21) dis with temperature depending on the sizes of may be reflected by the size of  $k_A$ , which may also be affected by the presence of some optimal sites on the nickel surface derivation of (21) but as demonstrated by Boudart (84, 87) this will still lead to rate (15) is assumed to be rate determining the appears. An expression of that form was functions being qualitatively correct. If step Surface heterogeneities were ignored in th derived by Lička (38).

Equation (21) may  $\sim$  power-rate law using the well-known approximation of the term ax/(1+ax) by  $^{1}$ 

$$= k_n p_{\text{C}_3\text{H}_4}^{1-2m} \cdot p_{\text{H}_4\text{O}^2(n-m)} p_{\text{H}_1}^{2(m-n)}, \quad (22)$$

coefficients obtained for catalyst A1 as shown in (13), and that the kinetic coefficient for steam may become less than 71 as observed for catalysts A35 and B4, but This expression may explain the kinetic

the reaction orders with respect to steam and hydrogen are not numerically equal as predicted.

cients to be deduced from Fig. 11 correlate with the kinetic order for steam indicating sated than accounted for in the simple In addition, the overall pressure coeffithe rôle of hydrogen to be more complisednence.

In Boudart's (34) recent kinetic interpres rium with nearby saturated sites on the metal surface different from the sites where tation of ethane hydrogenolysis hydrogen is assumed to be in a chemisorption equiliby he hydrocarbon reacts. If so, step (14)

ent activation energies for normal and fluenced the latter results, the relative re-This conclusion is supported by the differthough thermal pyrolysis must have inactivities appear to vary with temperature. cycloparaffins reported by Balashova, Slovokhotova and Balandin (40) (24)

(23)

 $H_2 + 2S_3 = 2(S_3) - H$ 

should be reformulated by:

 $G_{i}H_{i} + 2S_{1} + yS_{3} = (S_{1})_{2} - C_{2}H_{s}$ 

At 600°C Schnell (42) was able to detect substantial amounts of low olefins in re-

This implies that the term  $p_{c_2H_6}$  in (21)

 $+ y(S_3)-H + \frac{6-y-z}{2}H_2.$ 

(22) should be replaced by  $p_{c_2H_0}$ 

pH2 . Hence:

) pue

 $r = [1 + (2k'_A/k_r)(1/K_w) \cdot (p_{\text{G,H}}/p_{\text{H,0}}) \cdot p_{\text{H,1}}^{-(\nu/2)} + K_w(p_{\text{H,0}}/p_{\text{H,1}})^{2}]^{2})$  $k'_A(p_{\text{CiH}_a}/p_{\text{H}_1}^{\nu/2})$ 

$$= \frac{k_A(p_{\text{C,HJ}}/p_{\text{H}_1})''^2)}{(1 + (2k_A/k_r)(1/K_w) \cdot (p_{\text{C,HJ}}/p_{\text{H}_2})) \cdot p_{\text{H}_1}^{-1}(v'^2) + K_w(p_{\text{H}_2})/r}$$

(26) $p_{\rm H_1}^{2(m-n)-(n/2)y}$ .  $f = kp_{C_2H_s}^{1-2n} \cdot p_{H_2O^2(n-m)}$ 

reactivity trend of the higher hydrocarbons sorption simulating the steps suggested in hydrogenolysis remains speculative, and it may hardly be used for evaluation of the because the results from hydrogenolysis have been reported mainly for temperatures The simple sequence for hydrocarbon adbelow 400°C.

reactivity of branched hydrocarbons and cyclohexane deviates from the pattern generally observed in hydrogenolysis at tively the ends of chains (39). However, in reforming studies in the temperature range 250-350°C Balashova, Slovokhotova and tion of hydrocarbons and a much higher affins. At higher temperatures, 350-450°C, Phillips, Mulhall and Turner (7) observed partially decomposed hydrocarbons from drocarbons which were less reactive than normal paraffins. Methyl cyclohexane was found to be slightly less reactive than the present study at 500°C intermediates fere indicated only for benzene this being the least reactive hydrocarbon. In experi-Yoshitomi, Morita and Yamamoto (41) found cyclohexane to be more reactive than n-hexane above approx 800°C. Al-The apparent multiple fission of carboncarbon bonds and the relatively high low temperature, nickel attacking selec-Balandin (40) found a stepwise degradareactivity of n-paraffins than of cycloparthe reforming reaction with branched hynormal heptane and a small amount of ments at very high temperatures 750toluene was found among the products. . '⊃₀ġ

times. In the present experiments on butane sions and contact times (approx 2 × 10-3 tified in these studies performed at 400-525°C, the olefins observed by Schnell are reaction on the nickel surface but products from thermal pyrolysis or cracking on the carrier material, these reactions being more butane performed at very short contact sec) were within the range considered by most probably not intermediates from the by Schnell. On this basis, it appears reasonable to assume multiple fission of the peratures of 500°C and above, and to conreformers as being hydrogenated products reforming at atmospheric pressure conver-Schnell. Since no intermediates were idenpronounced at the temperature level applied carbon-carbon bonds on the nickel surface for most nonaromatic hydrocarbons at temsider minor amounts of ethane, propane and other higher hydrocarbons which might be observed in the effluent of some tubular forming experiments with from pyrolysis or cracking.

# 2. Catalyst Activity

nickel for reforming reactions, hydrotivity trends is observed. Another common The results shown in Table 7 demonstrated that the specific activity of fluenced by the carrier employed and by the presence of alkali. A parallelism of acfeature of these reactions is a resemblance of the trend of specific activities of different metals, ruthenium and rhodium being reported to show much higher specific activities for hydrogenolysis (11) and methanagenolysis and methanation is strongly in-

tion (43) than do nickel, platinum, and rier and the presence of alkali. This makes palladium. Contrary to this, ammonia decomposition appears unaffected by the cardoubtful the use of this reaction for activity tests of reforming catalysts as proposed by Merkel (44).

A carrier effect in reforming reactions nickel on silica. Balashova, Slovokhotova and Balandin explained the difference by a cyclohexane. However, as proven by the hydrogenolysis experiments, nickel on carbon may show poor activity in other reac-Slovokhotova and Balandin (4) who found nickel on carbon to be nearly inactive for reforming of cyclohexane compared with failing ability of coke for activation of steam since the two catalysts showed comparable activities for dehydrogenation of has also been described by Balashova, tions where steam is not involved.

A low activity for hydrogenolysis has based catalysts, the latter being the most curacy of the experiments does not allow dioxide Pour (45) found the specific activity of supported nickel to decrease in the effective. These results are in accordance based catalysts. For methanation of carbon also been reported by Sinfelt (11) for cobalt on carbon. Moreover, Sinfelt showed the specific activity for hydrogenolysis of silica-alumina-based nickel catalysts to be much less than those of alumina- or silicawith the trend observed for ethane reforming in the present study, although the aca distinction between alumina- and silicaorder chromia, alumina, silica.

mentioned by Andrew (3). The effect was by increasing the amount of sulfur, This forming reactions caused by alkali was explained by adsorption of alkali blocking When this explanation is applied for the over, it implies that alkali is adsorbed more specifically at the most active sites than is fall by increasing the amount of alkali than adsorption of alkali is not affecting the sites for ammonia decomposition. Moresulfur, because as mentioned earlier, the specific activity displays a more drastic The diminution of the activity for rethe most active sites on the nickel surface. present measuements, it implies that

behavior appears unlikely but, in princi it cannot be excluded, as demonstrated Roginskii (46).

Schoubye (12) reported low activities by assuming alkali to depress the affir observed a similar effect on a nickel all of the surface for hydrogen so that de and Taylor (47) observed a great effeg alkali on the kinetics of ethane hy genolysis on iron. This effect was explain drogenation of the adsorbed hydrocarb retarded. In the simple sequence (14), means a higher value of z. Shephard ina catalyst and noticed that remove The effect of alkali has also been alkali resulted in increased specific acti For methanation of carbon mond ported for hydrogenolysis. Cimino, Boi alkali-promoted catalysts.

position of ammonia appears "facile". The and methanation may therefore be charac-faized as structure-sensitive, while decomments and no correlation between surf was arrived at by Richardson (48). temperature range applied in the exi on a rôle of acid sites for hydrogeno The evaluation of interactions beti metal and support has been an intisubject of catalysis. Bifunctional cata is not very likely to be involved apart which cannot explain the different spe activities. As discussed earlier, cracking the carrier surface may be excluded in within the same range. A similar concil acidity and specific activity is appa the rôle of the carrier in steam adsor y-alumina show specific acti Thus catalysts based on magnesia,

ereas the activity showed no variation

methanation on nickel supported by mina Bousquet and Teichner (53) found influence on the specific activity of stallite size. The nickel crystallites of

crystallite size for nickel on silica.

rox 70 Å (17). In hydrogenolysis results ear ambiguous, as Sinfelt (11) found Feffect for nickel on silica-alumina,

> treaction rather than the collective properties of the bulk phase. Therefore, apply from extremely small metal crystallitics is used by Dalla Betta and Boudart (52) and the collection that can the carried intended. (49) and Szabo and Solymosi (50)atoms being important for the catall electronic interaction between the can doping of oxides supporting thin nic localized chemical properties of the surf change of the activation energy with films was assigned to a change of the se conductor properties of the carrier influemphasized in the work by Schwab metal phase. However, from later deve The electronic nature of the support ing the Fermi level of the electrons ments (51) there is much evidence for

y a result of crystallite size but rather a

action. Thus, Ponec and Sachtler (64)

deanwhile, surface heterogeneity is not tter of surface topography, with the stence of ensembles of nickel atoms on surface being optimal for the particular ction in question. The configuration of th ensembles may vary from reaction to ton their work on nickel-copper alloys gested that the ensembles required for

th crystallite size might be indicated for

catalyst with the smallest crystallites.

hough a decrease of the specific activity

outside the range of interest, and rening of ethane appears to be unaffected the crystallite size as shown in Fig. 4,

catalysts in the present study are nearly

bond or leading to isomerization contain a smaller number of adjacent nickel atoms reactions affecting the carbon-hydrogen genolytic splitting of the carbon-carbon than the ensembles required for hydrobond. the surface atoms of the metal is very likely. The measurements reported in le 4 and Fig. 3 are in accordance with statement, as the use of zirconia (D4) ing in low specific activities are accomhe presence of alkali (A34 and A35) re-

tion energies. Contrary to this, an eased activation energy observed when ng a nickel-copper catalyst (A13) may

ed by no detectable change of the ac-

s mentioned earlier the correlation wn in Fig. 6 indicates that different dific activities might be related to the

Explained by an influence on the nickel ace atoms of neighboring copper atoms. reforming reactions, hydrogenolysis

frogeneity of the nickel surface which sumably is influenced by the carrier. methanation may therefore be characdrystallite size because heterogeneity of the

ect of surface structure is often studied series of catalysts with varying metal ace is expected to increase by decreasthe crystallite size in the range below

anisotropy of the surface energy of the Unfortunately, very few studies have been contact angle for stabilized zirconia as cated by low contact angles that wetting several oxides. Contact angles on zirconia tion of impurities at the interface may facetting and other equilibrated shapes shapes of the crystals which could be stabilized by the support material. The interface is extremely complex as chemical reactions metal and by the interfacial energy (57). reported on this effect. For liquid nickel droplets at 1500°C Kingery (58, 59) found a significantly higher adhesion energy and correspondingly low interfacial energy and support than for alumina. Studies in vacuo of nickel droplets on various materials indiwas more pronounced on graphite than on and titania were lower than on magnesia, alumina and beryllia. Preferential adsorplower the interfacial energy and result in On larger crystals the number of special tion (55, 56) or the ensembles may be related to epitaxial relations or abnormal may be involved. In principle, the shape of ensembles may vary with crystal orientathe metal particle is influenced by (58, 60, 61).

the importance of various crystallite forms It is evident that these observations allow no conclusions to be made concerning the of the metal. In addition, use of data obtained for droplets at high temperature appears doubtful. However, there may be some indication that carrier materials showing high adhesion energy and thus influencing the shape of the metal particle cause a decrease of the number of ensemit is conceivable that the influence of alkali is caused by adsorption of alkali at the interface or on the metal changing the the planes. Shephard (33) also considered shape of the crystal or the facetting of for the activity for hydrogenolysis. Examieffect of the carrier on the catalytic activity bles for the reforming reactions. Moreover,

nation in the electron microscope of the catalysts of this study revealed no marked defined systems are required to evaluate differences between active and poor catalysts, and more detailed studies on wellthese effects.

## CONCLUSIONS

range by a simple power law, as the powers The kinetics and the specific activity for reforming reactions vary from catalyst to catalyst. The major differences in the kinetics are found in the influence of steam Active magnesia or the presence of alkali enhance steam adsorption. The kinetics partial pressure. This is related to the abilcannot be described in a broad temperature ity of the carrier material to adsorb steam. vary with temperature.

At temperatures around 500°C the results hydrocarbons and cycloparaffins appear face of the carbon-carbon bonds for most higher than those of normal paraffins indicate multiple fission on the nickel surof normal paraffins at equal partial pressures appears to increase with the number nonaromatic hydrocarbons. The reactivity of carbon atoms. The reactivity of branched whereas aromatics show poor reactivity.

same range for a great number of catalysts procedure. Use of supports such as zirconia activities, whereas some decrease of the alkali in various ways implies a significant The activities per unit nickel surface area, the specific activities, are within the port material, preparation or activation and carbon results in very poor specific specific activity is observed when using irrespective of the nickel surface area, supsilica-alumina and titania. Addition of drop in specific activity, the effect of potassium being larger than for sodium.

with results obtained or reported for ethane ing reactions are generally in accordance hydrogenolysis and methanation, whereas ammonia appears unaffected by the carrier and the presence of alkali. It is unlikely that this effect of the carrier and alkali is the specific activity for decomposition of related to the ability for steam adsorption The activity trends observed for reformor to surface acidity. Morever, electronic

interactions between carrier or alkalisand the metal may be excluded, since the effect the metal may be excluded, since the effect the metal may be excluded. activation energy.

The results indicate that the activity reforming correlates with the surface erogeneity as expressed by the adsorp adsorption of alkali on the interface of energies. This might affect the numbé of nitrogen on the nickel surface. It lieved that some carrier materials actions, hydrogenolysis and methanati metal may influence the surface stru by changes of the surface and inter ensembles being optimal for reformir

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